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Orals

O 1
Quick Annotator: an open source digital pathology tool for annotating objects 70 times faster than manual annotation
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Background: Machine learning approaches for the segmentation of histologic primitives (e.g., cell nuclei) in digital pathology (DP) Whole Slide Images (WSI) require large numbers of exemplars. Unfortunately, annotating each object is laborious and often intractable even in moderately sized cohorts. Here, we present an open source tool, Quick Annotator (QA), designed to improve the annotation efficiency of histologic primitives on WSIs of human annotators by 70× via the integration of deep learning (DL) and active learning. While the user annotates regions of interest (ROI) via an intuitive web interface, a DL model is concurrently optimized using these annotations and applied to the ROI. The user iteratively reviews DL results to either (a) bulk accept accurately annotated regions, or (b) correct erroneously segmented objects to improve subsequent model suggestions, before transitioning to other ROIs.

Methods: 3 Pancreatic adenocarcinoma (PAAD) TCGA WSIs were employed to evaluate the efficiency improvements afforded by QA. To estimate manual annotation speed, QuPath [1] was used to delineate cell nuclei in 2500×500 ROIs. QA usage time was recorded for annotating the equivalent of 600 500×500 ROIs, including the manual ROIs. Resulting annotations were quantitatively and qualitatively compared.

Results: QA employment yielded 345,722 nuclei in 535 min, or 10.8 nuclei per second (NPS). A similar feat employing QuPath necessitates ~685 h based on the 0.14 NPS performance witnessed in the 500×500 ROIs. Qualitatively, this 70× speed improvement does not appear to affect annotation quality (Fig. 2), which is reflected by a mean pixel-level f-score of 0.975 computed between manual and QA annotation results.

Conclusion: Our preliminary results suggest QA provides a 70× efficiency improvement while producing annotations highly concordant with manual efforts. The nature of the QA tool further suggests that it could be employed for annotation tasks involving other histologic primitives.

Fig. 2 | O 1 Fig. 2 shows (a, d) original 500×500 ROIs with (b, e) associated annotations overlaid in fuchsia. This manual result is compared with the output from QA (c, f), such that pixels in common between both results are white, and those missing from (b, e) appear green, and those unique to QA appear pink. Their high concordance, as indicated by the presence of mostly white pixels, is supported by f-scores of 0.99 (top) and 0.96 (bottom). Notably, although highly similar, QA’s annotations were produced over 70 times faster than manual efforts

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**Abstracts**

**O 2**

Prevalence of MET Exon 14 mutations or MET amplification in non-small cell lung cancer in Swiss patients

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**Background:** MET-targeted treatment is entering clinical practice for MET Exon 14-mutated or MET-amplified non-small cell lung cancer (NSCLC). MET exon 14 skipping mutations occur in 3% and MET amplification in 1 to 6% of unselected NSCLC. They are typically mutually exclusive with other oncogenic driver alterations. The aim of our study was to investigate the prevalence of these MET alterations in treatment-naïve pre-selected NSCLC from our routine clinical practice, were MET analyses were generally performed sequentially in NSCLC with high MET expression and wild type status for EGFR, KRAS, ALK and ROS1.

**Methods:** 580 consecutive NSCLC were retrospectively evaluated. High MET expression was defined as complete membranous MET staining with at least moderate intensity in ≥50% of tumor cells. MET exon 14 analysis was performed by Sanger sequencing. MET gene copy numbers were evaluated by fluorescence in situ hybridization (FISH). MET amplification was defined as a MET/CEN7 ratio of ≥2.0. MET immunohistochemistry (IHC) was performed on histology specimens and cellblocks, but not on conventional cytology preparations. Therefore, prevalence data of MET alterations were analyzed separately for IHC pre-selected and non-pre-selected NSCLC wild type for other oncogenic driver alterations.

**Results:** 66% (302/457) of NSCLC had a high MET expression. There was no difference between the prevalence of MET exon 14 mutations in IHC pre-selected and non-pre-selected NSCLC (8.3% (9/99) and 8.1% (7/86), respectively, p = 1.0). In contrast the prevalence of MET amplification was higher in MET IHC pre-selected NSCLC (11.5% (11/85) and 3.2% (1/30), respectively, p = 0.29).

**Conclusions:** MET exon 14 skipping mutations are enriched in NSCLC wild type for other oncogenic driver alterations with the prevalence of >8% being higher compared to unselected NSCLC (3%). High MET expression does not enrich for MET exon 14 mutations. However, MET IHC seems to be a cost-effective approach for pre-screening NSCLC for further MET FISH analysis.

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**O 4**

Two distinct immunopathological profiles in autopsy lungs of COVID-19

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**Coronavirus Disease 19 (COVID-19) is a respiratory disease caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), which has grown to a worldwide pandemic with substantial mortality. Immune mediated damage has been proposed as a pathogenic factor, but immune responses in lungs of COVID-19 patients remain poorly characterized. Therefore we conducted transcriptomic, histologic and cellular profiling of post mortem COVID-19 (n = 34 tissues from 16 patients) and normal lung tissues (n = 9 tissues from 6 patients).** Two distinct immunopathological reaction patterns of lethal COVID-19 were identified. One pattern showed high local expression of interferon stimulated genes (ISGhigh) and cytokines, high viral loads and limited pulmonary damage, the other pattern showed severely damaged lungs, low ISGs (ISGlow), low viral
loads and abundant infiltrating activated CD8+ T cells and macrophages. ISGhigh patients died significantly earlier after hospitalization than ISGlow patients. Our study may point to distinct stages of progression of COVID-19 lung disease and highlights the need for peripheral blood biomarkers that inform about patient lung status and guide treatment.

O 5
Development of a novel custom next generation sequencing panel to facilitate diagnosis and tumor classification of salivary gland neoplasms

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Background: The differential diagnosis and classification of salivary gland neoplasms can often be difficult and challenging. Recently, a range of highly specific molecular aberrations, in particular gene fusions as well as mutations, in different salivary gland tumor entities have been described. However, due to the frequent aberrations in otherwise rarely mutated genes, molecular testing is often expensive and time consuming as more than one test is required to cover all possible alterations. Therefore, we designed a novel RNA-based next generation sequencing panel including 27 genes, specifically involved in salivary gland and odontogenic neoplasms.

Methods: We designed a custom next generation sequencing (NGS) panel based on the Archer FusionPlex technology including 27 genes, known to be specifically involved in salivary gland and odontogenic neoplasms. Panel validation was performed on RNA from FFPE material of a variety of different salivary gland neoplasm entities and other tumors with known alterations. The results were compared to the results of our current diagnostic standard for the respective alterations (FISH, NGS, Sanger sequencing).

Results: Our customized NGS panel reliably detected all known translocations and mutations using RNA from FFPE tissue. The panel allowed analysis of common alterations in salivary gland tumors including the detection of NTRK1–3 fusions that can be used as therapeutic targets. Furthermore, the panel could successfully be used in one case of a difficult-to-classify adenoid–cystic carcinoma mimicking pleomorphic adenoma by detection of a MYBL1–NFI B fusion.

Conclusion: This newly developed, salivary gland neoplasm specific NGS panel reliably detects gene fusions and mutations known to be associated with specific entities. This approach can be a comprehensive, time-efficient and powerful tool to facilitate the diagnosis and classification in difficult cases.

Posters

P 1
Retrospective post-mortem SARS-CoV-2 RT-PCR of autopsies with COVID-19 suggestive pathology supports absence of lethal community spread in Basel, Switzerland before February 2020

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Background: The first case of Coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) was recorded in Hubei in December 2019. However, the extent of early community spread in Central Europe before this period is unknown.

A high proportion of asymptomatic cases and undocumented infections, high transmissibility and phylogenetic genomic diversity have engendered the controversial possibility of early international community spread of SARS-CoV-2 before its emergence in China.

Methods: To assess the early presence of lethal COVID-19 in Switzerland, a retrospective analysis of deaths at University Hospital Basel between October 2019 and February 2020 (n = 310) was performed, comparing the incidence of clinical causes of death with March 2020 (n = 72), the month during which the first lethal COVID-19 cases were reported. Trends of COVID-19 suggestive sequelae, such as bronchopneumonia with organ embolisms (PE) were evaluated. In cases where an autopsy was performed (n = 71), analogous analyses were conducted on the cause of death and pulmonary histological findings (Fig. 1). Eight cases with COVID-19 suggestive clinical history and histopathology between October 2019 and February 2020, and three cases before October 2019 were selected for SARS-CoV-2 RT-PCR.
Results: A statistically significant rise in pulmonary causes of death were observed in March 2020 (p = 0.03), consistent with the reported emergence of lethal COVID-19 in Switzerland (Fig. 2). A rise in lethal bronchopneumonia was observed between December 2019 and February 2020, which was likely seasonal. The incidence of lethal ARDS and PE was uniformly low between October 2019 and February 2020. All autopsy cases analyzed by means of SARS-CoV-2 RT-PCR yielded negative results (Fig. 3).

Conclusion: Our data suggests the absence of early lethal community spread of COVID-19 in Basel before its initial reported emergence in Switzerland in March 2020.

Fig. 1 | P 2 ▲ Clinical Characteristics of Patients with positive and negative myocardial SARS-CoV-2 RT-PCR compared to controls * Significance testing between 3 groups (RT-PCR positive, negative and controls): categorical – Pearson’s Chi Square Test for Independence; parametric, continuous, -one-way ANOVA; ordinal/non-parametric – Kruskall Wallis one-way ANOVA. p-Values considered significant at the 0.05 level

** Significance testing between 2 groups (COVID-19 patients and controls): categorical – Pearson’s Chi Square Test for Independence; parametric, continuous, – independent samples t-test; ordinal/non-parametric – Mann Whitney U test. p-Values considered significant at the 0.05 level.

*** Refer to Materials and Methods for diagnostic criteria of myocarditis

Fig. 2 | P 2 ▲ Pathology Findings of Patients with positive and negative SARS-CoV-2 RT-PCR compared to controls

* Significance testing between 3 groups (RT-PCR positive, negative and controls): categorical – Pearson’s Chi Square Test for Independence; parametric, continuous, -one-way ANOVA; ordinal/non-parametric – Kruskall Wallis one-way ANOVA. p-Values considered significant at the 0.05 level

** Significance testing between 2 groups (COVID-19 patients and controls): categorical – Chi Square; parametric, continuous – independent samples t-test; ordinal/non-parametric – Mann Whitney U test. p-Values considered significant at the 0.05 level

*** Refer to Materials and Methods for diagnostic criteria of myocarditis
Results: COVID-19 patients had a higher mean age (76 ± 13 vs. 64 ± 14 years, p = 0.04) and increased incidence of hypertension (91 vs. 70 %, p < 0.01), coronary artery disease (61 vs. 20 %, p = 0.03) and diabetes mellitus (48 vs. 10 %, p = 0.04) than controls (Fig. 1). COVID-19 patients with a positive RT-PCR result died earlier after hospital admission (5 ± 3 vs. 14 ± 8 days, p < 0.01). Increased capillary dilation, capillary fibrin and microhemorrhage were observed in RT-PCR positive myocardium (Fig. 2); Spearman's Rho analysis revealed a positive correlation amongst these factors (Fig. 3). All cases fulfilling diagnostic criteria for myocarditis (borderline n = 4, lymphohistiocytic n = 1) were RT-PCR negative. In 3 cases, in-situ hybridization revealed viral RNA in interstitial cells.

Conclusions: Myocardial capillary dilation, fibrin deposition and microhemorrhage may be the histomorphological correlate of the hypercoagulable state observed in critically-ill COVID-19 patients and are thus suggestive of a generalized vasculopathic component in its pathophysiology. RT-PCR negativity in cases with myocarditis implies a secondary immunological response, which warrants further characterization.

Methods: Autopsies of 23 patients with COVID-19 were performed at University Hospital Basel and Cantonal Hospital Baselland, Liestal. Representative myocardial tissue sections were collected. Based on Ct-values of myocardial RT-PCR, cases were categorized as SARS-CoV-2 positive (n = 15) and negative (n = 8). Autopsies of patients without COVID-19 (n = 10) with similar clinical sequelae (diffuse alveolar damage, pulmonary thromboembolism, bronchopneumonia) were selected as controls. Histological analysis was performed on H&E and CAB sections and with immunohistochemistry (CD3, CD20, CD68, Fibrin, ACE2, SARS-CoV-2 N-Protein). Histological characteristics were scored by ordinal and/or categorical grading. The presence of myocarditis was evaluated according to Dallas criteria. Five RT-PCR positive cases underwent in-situ hybridization of viral S- and ORF1ab RNA. In two cases, electron microscopy was performed.

Results: SARS-CoV-2 virus was detected in formalin-fixed paraffin-embedded (FFPE) tissue samples, both from autopsies and from surgical and biopptic specimens. We describe the methodology set up at the University-Institute of Pathology in Lausanne.

Methods: Total RNA was extracted from FFPE samples from lung, liver and heart of 12 patients autopsied with intra-vitam confirmed COVID-19. One-step reverse transcriptase quantitative PCR (RT-qPCR) was performed for SARS-CoV-2 mRNAs (E and RdRP genes) as well as for human M-STN as internal control of RNA quality, using commercially available assays (Roche). Specific in vitro-transcribed RNA standards were used in each experiment for determination of viral copy numbers.

Results: Optimal RNA input was 50–250 pg per reaction. Limit of detection of E gene and RdRP was 7 and 27 copies per reaction, respectively (as determined by Probit analysis). SARS-CoV-2 was detected in 12/12 lung, 6/11 liver and 6/11 heart FFPE samples (one liver sample and one heart sample had an inadequate internal control), with very variable virus levels. The highest amounts were detected in the lung (range E gene: 11–537,000 copies per reaction; RdRP: 0–64,345), followed by heart (E gene: 0–9756; RdRP: 0–129) and liver (E gene: 0–583; RdRP: 0–9). All samples found positive for E gene were also positive for RdRP, but not reciprocally.

Conclusions: We show that a commercially available kit can be used for sensitive and quantitative SARS-CoV-2 virus detection in FFPE tissues. The limit of detection was lower for E gene than for RdRP. A suitable internal control for RNA quality is highly recommended in order to rule out false negative results.

P 4 Mutational landscape of marginal zone B-cell lymphomas of various origin: organotypic pathways and diagnostic potential for organ assignation

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Background: Marginal zone lymphomas (MZL) comprise nodal (NMZL), splenic (SMZL), and extranodal (EMZL) instances, accounting for 10, 15 and 75 %, respectively, and, as a whole, 12 % of all B-cell neoplasms. MZL have no disease-defining phenotype and the diagnostic borders to other lymphomas are blurred. MZL are known to bear typical site-of-origin translocations, yet their mutational landscape has not been integratively analyzed.

Principals/Methodology: Gene mutation data of 916 MZL, namely 287 ocular adnexal (OA), 93 pulmonary (P), 71 salivary gland (SG), 59 gastric (G), 38 cutaneous, 18 thyroid (T), 15 dura matter (D), 115 NMZL and 220 SMZL were retrieved from our own digital archives and published other groups’ studies. Data were pooled and used for inter-entity comparison. Number and frequency of mutated cases (≥2 and/or ≥5 %/entity), were set to generate a gene shortlist. Statistical differences were calculated by the two-tailed Fisher’s exact test; P-values were corrected for multiple testing (p < 0.005).

Results: Out of 223 genes, mutational data in 11 appeared to be of importance respecting MZL origin. NMZL more commonly displayed mutations...
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in KMT2D, FAS, TNFRSF14 or EZH2 compared to SMZL, in KMT2D or TNFRSF14–compared to OAMZL, and in KMT2D only–compared to SGMZL. Mutations in NFI were conjoined to GMZL compared to PMZL and DMZL. TMZL more often harbored mutations in TET2 compared to PMZL, SGMZL and NMZL. In PMZL, mutations in KMT2D or B2M were more frequent than in OAMZL. OAMZL more commonly harbored mutations in TNFAIP3 than NMZL and GMZL. Finally, in DMZL mutations in TNFAIP3 were more common than in SMZL.

Conclusion: Mutations of distinct genes show origin-preferential distribution among MZL. Recognition of such mutational distribution patterns may be of potential help assigning MZL origin in difficult cases and possibly also pave the way for novel tailored treatment concepts.

P 5
Quality comparison of computer-assisted T-cell and tumor-bud detection in pT1 colorectal cancer from four different pathology institutions*

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Background: Tumor budding and T-cell infiltrates are robust prognostic biomarkers in colorectal cancer. The identification of tumor buds and T-cells can be expedited and facilitated using deep learning. However, due to biomarkers in colorectal cancer. The identification of tumor buds and T-cells is more prone to pre-analytical factors. A variety of approaches can be used to address these issues in the development and validation of automated detection methods.

Methods: We validated deep learning algorithms for tumor budding and CD8+ T-cell detection on double-stained immunohistochemical slides (AE1-AE3 and CD8) from our pT1 colorectal cancer cohort from 4 different institutions, originating from 1992–2017. At least 20% of slides from each institute (total 76) were randomly selected for quality control. Staining and scanning was performed in a single laboratory using the same protocol. For comparison, a single observer identified tumor buds within the tumor budding hotspot (0.785 mm²) as well as correcting the T-cell algorithm output.

Results: Fig. 1 shows the number of correctly identified tumor buds, Fig. 2 the T-cell detection. The most consistent performance is achieved on slides from Institute A. Notably, institution C, which achieves the best results for T-cell detection shows the worst results for tumor bud detection. Upon review, heterogeneity in staining intensity between slides and within the same slide emerged as a major factor for tumor buds missed by the algorithm.

Conclusion: The algorithm for T-cell detection achieves better results for all institutions in comparison to tumor bud detection. This suggests that different algorithms face unique challenges in detecting individual objects. T-cell detection appears relatively robust, whereas in addition to known challenges in high inter- and intra-observer variability, tumor bud detection is more prone to pre-analytical factors. A variety of approaches can be used to address these issues in the development and validation of automated detection methods.

* Student paper

P 6
PatchSorter: a high throughput digital pathology tool for cell labeling

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Background: Recently significant interest in development of computational pathology tools for diagnosis, prognosis, and therapy response of many diseases has emerged. These approaches typically rely on accurate understanding of both location and cell type present within histology slide images. While image analysis algorithms can identify the location of millions of cells on digitized pathology images, a subsequent step is needed to identify cell type. Manually assigning labels (e.g., inflammatory, epithelial) to each cell individually remains an intractable task. Here, we present PatchSorter, a browser-based high-throughput cell labeling tool which enables the user to review and assign labels at a group, as opposed to individual cell level, thus greatly improving labeling efficiency. As the user classifies groups of cells, PS’s deep learning component iteratively increases separation between target cell groups within a low dimensional representational space (Fig. 1), thus further improving labeling efficiency.

Methods: A random subset of 14,330 cells, with ground truth labels indicating membership in either inflammatory or epithelial classes, from the multi-organ 27 H&E image Hover-net dataset [1] was used to evaluate PS. To estimate manual labeling speed, a Photoshop clone (Gimp) was used to label 120 cells from a representative image. Labels generated from PS were quantitatively compared against provided inflammatory or epithelial class labels using the accuracy metric.

Results: Labeling 14,330 cells using PS required 9607 s, yielding 1.5 labels per second (LPS). The estimated time to perform the same task manually would take 23,883 s, based on the .6 LPS recorded when manually using Gimp. Qualitatively, this 60% speed improvement only modestly affects labeling fidelity, as indicated by a 93% accuracy between PS produced labels and ground truth.

Conclusion: PatchSorter provides a 60% efficiency improvement for assigning cells to appropriate cell-type, while also yielding a 93% accuracy. Additional, independent validation and user testing is warranted.
Fig. 1 [P 6] (Top) To begin, the user lassos points of interest in the Cartesian plot (left) which are subsequently presented for review and bulk labeling (right). (Bottom) As labeling progresses, the feature space is improved, further purifying and separating predicted groups in the plot to facilitate improved efficiency in labeling.

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P 7
Mutational profiles of primary pulmonary adenocarcinomas and paired brain metastases
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Background: Brain metastases (BM) are frequent and clinically relevant in lung cancer patients. Molecular data on paired BM and primary lung cancer is scarce.

Methods: Consecutive solid cancer BM diagnosed at the Institute of Pathology Bern from 2000 to 2015 were retrospectively assembled. Paired tissue from pulmonary adenocarcinomas and BM was analyzed by next generation sequencing (NGS; oncomine comprehensive cancer panel v3). Results were compared to routine diagnostics and the TCGA dataset.

Results: NGS was successful in 54/57 consecutive tumor pairs accessible for molecular analysis. Mutation frequencies were in the range expected from TCGA. Primary tumors showed most frequently TP53 missense and truncating mutations (33/57, 60%), activating KRAS mutations and amplifications (32/57, 58%), MYC mutations and amplifications (9/57, 16%) and STK11 missense and truncating mutations (9/57, 16%). Significantly more KRAS mutations were detected at primary sites compared to in-house data from routine diagnostics and the TCGA (KRAS G13C mutation; p < 0.02). Most genetic alterations were preserved in the BM. Oncoprint analyses revealed a bias towards the EGFR signaling pathway in alterations private to BM, notably, secondary KRAS (22%), NFI (9%), MET (6%) and MYC alterations (13%).

Five models of progression emerged from the genetic alterations: Synchronous BM (10/57; 19%), showed the same set of mutations as the primary sites (P = BM). Private alterations were seen in 12/57 (22%) primary tumors (P > BM, most frequently RICTOR) and 14/57 (26%) BM, without enrichment of genes (P < BM). 14/57 (26%) patients had shared and private mutations at both sites (PBM). Frequent primary BM mutations were additional KRAS mutations/copy number gains (21% 21%). Only one case showed private KRAS alterations in the primary tumor.

4/57 (7%) tumor pairs had no shared mutations (P/BM), with different variants of KRAS and TP53.

Conclusions: Our data point to an important role of KRAS in the pathobiology of BM.

P 8
Putative malignant pleural mesothelioma in situ (MPMIS) with sequential acquisition of genomic alterations on fluorescence in situ hybridization (FISH) examination
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Background: The existence of a malignant mesothelioma (MPM) in situ has long been postulated. Here, we present a case of a 57-year-old man with initial presentation as a putative MPMIS evolving to a diffuse malignant pleural mesothelioma (MPM) during an interval of 8 months. Repeated FISH examination on pleural effusion from both lesions identified sequential acquisition of genomic alterations. Interestingly, in the initial pleural effusion, a substantial number of the mesothelial cells were encircled by lymphocytes, which we refer to as satellitosis.

Methods: Immunohistochemistry (IC) as well as multiprobe FISH assay were performed on the cytological specimens of both, the putative MPMIS as well as on the diffuse MPM. Further, in order to determine the incidence of satellitosis of mesothelial cells by lymphocytes, we reviewed 60 cases from our archive with proven MPM and 60 cases with the diagnosis of “reactive mesothelial cells”.

Results: IC analysis–loss of BAP1 and positivity for calretinin–confirmed the mesothelial and neoplastic nature in both specimens (Fig. 1). Further, FISH analysis showed homozygous deletion of 9p21 in the initial specimen and unbalanced polysomy of the chromosomes 3, 7, and 17, in addition to the loss of 9p21 in the second specimen (Fig. 1). In the cases reviewed, “satellitosis” was identified in a total of 5 cases, 2 cases with MPM, and 3 cases with reactive mesothelial cells (Fig. 2).

Conclusions: FISH analysis can serve as a screening tool to detect MPMIS in patients presenting with no clinical evidence of mesothelioma. Genomic transition from a diploid to an aneuploid state might play a role in progression from MPMIS to invasive MPM. Satellitosis of mesothelial cells by lymphocytes in pleural effusion cytology is a rare and not tumor-specific phenomenon with unclear relevance.
Abstracts

P 9
Progressive disease in sentinel-negative melanoma patients: Biological differences and importance of sentinel lymph node biopsy

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Background: Among the most important prognostic factors in melanoma is the sentinel lymph node (SLN) status.

Methods: Using our electronic database we identified 109 of 890 SLN-negative patients with progressive disease (PD). These patients were characterized for melanoma type, molecular type, sequence and extent of metastatic spread.

Results: A total of 61 of 109 SLN-negative patients had PD in the SLN basin indicating false-negative SLN (group-1). 48 of 109 patients had PD at distant sites and were therefore impossible to be identified using SLN biopsy (group-2). Despite distant spread these patients had significantly more single organ metastasis (p < 0.001) and significantly longer disease-free survival (p = 0.001) compared to group-1. Additionally, to significant differences on a molecular basis between the two groups (p = 0.01), all lentigo maligna and spindle-cell-melanomas belonged to group-2 and all, except one lentigo maligna melanoma, had single visceral metastasis.

Conclusion: Two different biological groups among SLN-negative patients with PD were demonstrated. Extravascular migratory-metastasis, rather than hematogenous spread, might be responsible for the observed PD with single organ involvement.

Fig. 1 | P 9  ▶

Fig. 1 | P 8  ▶

Fig. 2 | P 8  ▶

Fig. 2 | P 9  ▶

Fig. 1 | P 9  ▶
Background: Diffuse pulmonary meningotheliomatosis (DPM) is a rare condition. We report a case with review of the literature.

Methods: 72-year old woman with rheumatoid arthritis, exertional dyspnea and multiple bilateral centrilobular distributed noduli underwent surgical biopsy in order to prove an interstitial lung disease. Immunohistochemical studies and Next Generation Sequencing were performed.

Results: Histology revealed lung tissue with multiple small nodules of epithelial cells with round to oval nuclei devoid of atypia and eosinophilic cytoplasm, mainly distributed in perivenular location and along the walls of alveolar septae. The nodules measured up to 5 mm in greatest dimension and demarcated sharply from the adjacent pulmonary parenchyma. Immunohistochemical studies showed a positivity of the tumor cells for epithelial membrane antigen, progesterone receptor and CD56. There was a negative staining for smooth muscle actin, CD34, CD117, S100 HMB45, AE1/3, Synaptophysin and Chromogranin A. The proliferation marker Ki-67 was 2%.

Next Generation Sequencing showed gene mutations of NOTCH2, MSH2 and SETD2.

Conclusions: DPM is a rare entity which can clinically and histologically lead to misinterpretation of other diseases. Initially described as chemodectomas, subsequent ultrastructural and immunohistochemical studies demonstrated the characteristics of meningothelial cells. Our immunohistochemical results confirm previous reports of meningothelial differentiation in DPM.

To our knowledge this is the first case of DPM described in the context of rheumatoid arthritis and gene mutations of NOTCH2, MSH2 and SETD2. We discuss the controversial nature of DPM.

P 11
Impact of extracellular mucin detection in colorectal cancer using a novel unsupervised deep learning method

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Background: Mucinous colorectal cancers are defined as containing >50% extracellular mucin and are found in approximately 12% of all cases. This cut-off is, however, arbitrary and the clinical relevance of mucin is controversial. The aim of this study was to develop a deep learning algorithm...
for quantification of the percentage of mucin within the tumor area and to determine the association of mucin with clinicopathological features.

**Methods:** We scanned 904 slides from 452 cases. Histological subtype and clinicopathological features were extracted from reports. We developed an unsupervised segmentation method to detect the percentage of extracellular mucin/tumor area by assigning a semantic label to each pixel in images and classifying them. The proposed method, namely Group Affinity Unsupervised (GAUS) learning has three main steps (Fig. 1): 1) Forward process to create an output image based on the classification of CNN visual features 2) Creation of a target image from the similarity group matching of color, spatial coordinate and prior tissue pattern 3) Backpropagation process with a loss function to train and update the network parameters. Inter-observer agreement between algorithm and pathologists’ scores was tested on 163 slides.

**Results:** Fig. 2 shows an example of segmentation output. Inter-observer agreement between pathologists and algorithm was excellent (Fig. 3; ICC = 0.917). Mucinous histology (>50 % mucin) was associated with larger tumor size (p = 0.008), right-sided tumor location (p = 0.0428), more advanced pT1 (p = 0.002), less venous invasion (p = 0.0451), less tumor budding (p = 0.023) and high levels of microsatellite instability (MSI-H) (p < 0.0001). The presence of any intratumoral mucin (>0 %) correlated with BRAF mutation (p = 0.0329).

**Conclusions:** We introduced an effective method for the mucin quantification in colorectal cancers, taking advantage of unsupervised learning without training data or pre-trained network. Moreover, tumors with any extracellular mucin behave similarly to cancers with mucinous histology (>50 % mucin). A more comprehensive investigation into the topic of mucin in colorectal cancer is warranted.

**P 12**

**Postmortem pulmonary pathology and virus detection in patients with COVID-19 infection: single-center report of 12 consecutive autopsies from Lausanne, Switzerland**

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**Background:** The novel severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) pandemic has caused >850,000 deaths worldwide, with a disproportionate lack of autopsy studies.

**Methods:** We report postmortem pulmonary findings focusing on histology and virus detection in 12 of 14 consecutive patients who died with confirmed COVID-19. Two patients declined participation in research. SARS-CoV-2 virus detection in formalin-fixed paraffin embedded tissue was performed by reverse transcriptase quantitative PCR (RT-qPCR) and in-situ RNA hybridization (RNAscope).

**Results:** The 5 women and 7 men (median age: 73 years; range 36–96) died 3–38 days after onset of symptoms (median: 14.5 days). Eight patients received mechanical ventilation for 2–19 days. Eleven patients showed diffuse alveolar damage (DAD) in different stages of evolution: 1 in early exudative phase, with congestion, edema and hyaline membrane formation; 6 (50 %) in late exudative phase, with additional thickening of alveolar septa and interstitial lympho-plasmacellular infiltrates; 4 (33 %) as proliferative phase/organizing DAD, in one patient presenting as acute fibroinflammatory and organizing pneumonia. Acute broncho-pneumonia was present in 6 patients (50 %), in one case without underlying DAD. One patient showed invasive aspergillosis with necrotic bronchiitis. Microthrombi were present only in exudative DAD in 5 patients. True necrotizing vasculitis was absent. RT-qPCR detected high amounts of virus in 6 patients (50 %) with exudative DAD and symptom-duration ≤14 days, supported by in-situ visualization of viral RNA in 3/3 cases analyzed. The 6 patients with borderline low viral copy levels were symptomatic for ≥15 days, comprising all 4 cases with organizing DAD, the patient without DAD and one case of exudative DAD.

**Conclusions:** We show the high prevalence of DAD as a reaction pattern in COVID-19, the high number of overlying acute bronchopneumonia, and convincing pulmonary virus detection limited to patients who died ≤2 weeks after onset of symptoms, correlating with pre-organization phases of DAD.

**P 13**

**Metastases in hepatocellular carcinoma retain morphology but show difference in methylation profile**

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**Background:** Metastasis is the leading cause of death in cancer patients. Especially in hepatocellular carcinoma (HCC), the prognosis is poor once metastases have formed. Little is known about the mechanisms of metastasis formation in liver cancer and about the changes that occur from the primary tumor to metastasis. The aim of our study was therefore to gain insight into the similarities and differences between primary tumor and metastasis in patients with hepatocellular carcinoma.

**Methods:** We analyzed morphology between primary tumor (n = 24) and matched metastasis present in various organs (e.g. lung, lymph node, or adrenal glands). In particular, we compared histological growth patterns (e.g. trabecular, solid or pseudoglandular), cytological characteristics, and differentiation grade. In addition, we compared methylation profile of several primary HCCs and their matched metastasis.
Results: 20 out of 24 patients (83.3 %) showed identical morphology between primary tumor and matched metastasis. In four cases (16.6 %), the metastasis did not show the same morphology as the primary tumor, for example the metastasis contained only one histological growth pattern whereas the primary tumor contained various growth patterns. In contrast, methylation analysis displayed a high degree of intra-donor clonality and exhibited patterns of divergent evolution. In particular, metastasis-associated fibroblasts showed the greatest extent of epigenetic reprogramming.

Conclusions: Our study revealed a high concordance of morphology but pronounced differences in methylation profile between primary and matched metastasis of HCC patients.

P 14
Posttransplant EBV-negative diffuse large B-cell lymphoma with microsatellite instability: an unexpected finding revealed by NGS
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Background: Microsatellite instability (MSI) is very rare in lymphoid neoplasms, but may occur in immunodeficiency-related lymphomas such as posttransplant lymphoproliferative disorders (PTLD). A compromised immunity might indeed fail to counter the development of hypermutated tumours, despite numerous neoantigens. Moreover, some immunosuppressive treatments like azathioprine are thought to induce epigenetic silencing of mismatch repair (MMR) enzymes. We present a case of PTLD with MSI revealed fortuitously by NGS.

Methods: A customized 54-gene-NGS panel, designed for B-cell lymphomas and covering a total of 134 kb, was applied to a PTLD recurrence. A mutational signature was determined based on the mutation patterns. MSI status and MMR proteins were further explored by PCR and immunohistochemistry (IHC).

Results: A 1983-born man underwent renal transplantation in 1998 for post-meningococcemic renal failure, pancreatic allograft in 2011 for type 1 diabetes, and suffers from a severe Crohn disease since 2013. He has received several immunosuppressive treatments including corticoids, calcineurin inhibitors, mycophenolate and azathioprine. In August 2019, gastric and pulmonary biopsies revealed an EBV-negative diffuse large B-cell lymphoma-type PTLD. Despite chemotherapy, a gastric PTLD recurrence was diagnosed in January 2020. NGS, performed to search for potential treatment targets, identified numerous non-synonymous mutations (likely equivalent to a tumour mutation burden >100/Mb), mostly transitions and small indels in homopolymeric sequences, suggestive of an MSI-associated mutational signature. MMR IHC showed a complete loss of MSH2 expression and a partial loss of MSH6. PCR confirmed MSI (5 instable markers/5).

Conclusions: Although small NGS panels are not considered suitable to determine a mutational signature, particular mutation patterns may hint towards a specific etiopathogenesis such as MSI, opening up new treatment opportunities. Reduction of immunosuppression and/or immune checkpoint inhibitors could theoretically have restored immune recognition of this hypermutated PTLD, but were contraindicated because of the patient’s comorbidities. He is currently responding to polatuzumab (anti-CD79b)–bendamustine.

P 15
Automatic detection of h.pylori in gastric biopsies*
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Background: Infection with Helicobacter pylori is commonly diagnosed in Giemsa-stained slides from gastric biopsies. This diagnostic task occurs daily and can be challenging and time-consuming. Here, we sought to au-
tomate the detection of *H. pylori* in whole slide images of Giemsa-stained gastric biopsies using deep learning methods.

**Methods:** 41 slides from different patients were scanned using a Hamamatsu S360 slide scanner at 40× magnification (resolution: 0.23 µm/pixel). The lumen region, where *H. pylori* is commonly found, was initially segmented using a U-Net convolutional neural network (CNN), which had been trained with 1048 annotated images. Subsequently, another CNN assessed the presence of *H. pylori* in patches extracted according to one of two strategies we evaluated: A) Laplacian of Gaussian blob detection was used to identify patches containing objects of the same size as *H. pylori* and B) patches were extracted directly from the lumen region using a grid (Fig. 1). We compared the three state-of-the-art CNN architectures MobileNetV2, ResNet50V2, and Xception trained on a dataset of 754 images, created by an expert pathologist.

**Results:** 98 % of lumen regions were detected by the U-Net, the dice overlap index was 0.89 ± 0.07 (Fig. 2), and the search space was narrowed down to 3.7 ± 2.3 %. 99 % of all bacteria were annotated as candidates by blob detection. The best performing architectures were Xception in strategy B (F1 = 0.989 ± 0.009), and ResNet50V2 (F1 = 0.859 ± 0.053) in strategy A. The assembled pipeline with strategy A/B detected 26 %/35 % true-positive, 25 %/37 % false-positive, and 49 %/28 % objects that could not be classified by our pathologist.

**Conclusions:** Despite the high number of false-positive results, we succeeded in demonstrating that it is possible to automatically detect *H. pylori* in gastric biopsies. Our results may improve with a larger database of annotated *H. pylori* images and higher-resolution scans.

* Student paper

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**Abstracts**

**P 16**

Superior survival prediction using a combined score of major pathological response, ypT and ypN compared to the standard TNM model in non-small cell lung cancer after neoadjuvant therapy*

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**Background:** The current standard prognostic models in non-small cell lung cancer patients after neoadjuvant treatment are the TNM classification and major pathological response (MPR) with a cut-off of 10 % residual tumor. We aimed to integrate tumor regression in a prognostic score (PRSC) to improve the possibility of clinical decision making based on a TNM and morphology-based system with increased predictive power.

**Methods:** 117 consecutive patients resected after neoadjuvant therapy between 2000 and 2016 were included in this monocentric retrospective study. During histopathological re-evaluation, the amount of residual tumor in the primary tumor bed was assessed according to the recommendations of the IASLC and tumor regression was graded into five categories: 0 %, <1 %, 1–10 %, 11–49 % and ≥50 % residual tumor. After confirming the prognostic significance of the TNM staging system and tumor regression, we included ypT, ypN and major pathological response (MPR) in a PRSC generating a three-tier classification. Cox-proportional hazard models were used for univariate and multivariate analyses. The Akaike and Bayesian Information Criterion were used to compare the goodness-of-fit.

**Results:** The isolated ypT-categories (p < 0.001, p < 0.001) and combined TNM8 stages (p = 0.004, p = 0.013) had significant impact on overall (OS) and disease-free survival (DFS) stratification. We confirmed the importance of histology-specific interpretation of MPR confirming the cut-offs at 65 and 10 % residual tumor in adenocarcinoma and non-adenocarcinoma, respectively, as previously reported. MPR significantly stratified OS (p = 0.01) and DFS (p < 0.001). The PRSC significantly stratified a low-, in-

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**Fig. 2 | P 15**
Intermediate- and high-risk group ($p < 0.001$ for OS/DFS). The PRSC yielded a better fitting model compared to the TNM classification or MPR alone. **Conclusion:** We established a prognostic score with the potential to more accurately predict patient survival, which is readily applicable in routine diagnostics.

* Student paper

**P 17**

**Significance of tumor regression in lymph node metastases of esophageal adenocarcinomas after neoadjuvant therapy**

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**Background:** Tumor regression following neoadjuvant treatment can be observed in lymph node (LN) metastases similar to the primary tumor in esophageal adenocarcinomas (EAC). We evaluated the prognostic significance of tumor regression in LN metastases locally advanced EAC following neoadjuvant treatment.

**Methods:** 239 EAC patients treated with neoadjuvant radiochemotherapy (RCTX) or chemotherapy (CTX) followed by esophagectomy were analyzed. We examined retrospectively the LN for histopathologic signs of regression, i.e. nodular fibrosis and acellular mucin. LN classification performed according to two parameters: presence (−) or absence (+) of residual tumor and regression characteristics in the LN, resulting in four categories: LN−/REG−, LN−/REG+, LN+/REG+, LN+/REG−. **Results:** LN metastases with residual tumor were detectable in 117/239 (49%) cases. Regression in LN were observed in 85/239 cases (35.5%). The distribution of the LN/REG categories were as follows: 97 patients (40.6%) were LN−/REG−, 25 patients (10.5%) were LN−/REG+, 60 patients (25.1%) were LN+/REG+ and 57 patients (23.8%) LN+/REG−. The LN/Reg categorization had a significant prognostic value in univariate analysis ($p < 0.001$) and multivariate analysis (HR = 1.326, $p = 0.002$) with similar results for the subgroups of patients treated with RCTX or CTX. The prognosis of LN−/REG+ was worse than LN−/REG− but better than both LN+ categories, which was clearly seen in the Kaplan Meier curves but did not reach statistical significance ($p = 0.104$ and $p = 0.090$, respectively). In contrast, there was no difference between LN+/REG+ and LN+/REG− ($p = 0.802$).

**Conclusions:** Regression in LN metastases of EAC can be observed in a significant number of patients after neoadjuvant therapy. The most relevant negative prognostic factor is the presence of LN metastases, independent of the degree of regression. Complete regression of former LN metastases in comparison to ‘true’ negative LN seems to be of prognostic relevance but additional studies are needed to confirm this trend seen in our study.

* Student paper

**P 18**

**Immunohistochemistry for SARS-CoV-2 in autopic pulmonary tissue**

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**Introduction:** Covid-19 is primarily a disease of the lungs. SARS-CoV-2, the virus causing the disease, is usually detected by PCR-based methods. In contrast, little is known about the value of immunohistochemistry (IHC) for visualizing SARS-CoV-2 in the tissue.

**Materials and methods:** Pulmonary FFPE autopsy tissue of patients with confirmed Covid-19 infection was analyzed by IHC using an anti-SARS-CoV nucleocapsid (N) antibody. In addition, the copy number of the viral RNA was determined by quantitative RT-PCR, which targets three genomic regions specific for SARS-CoV-2 and the human control gene RNaseP.

**Results:** 88 pulmonary samples of 19 Covid-19 patients were analyzed. IHC was positive in 63 (71.6%) samples of 10 (55.6%) patients and RNA detected in 83 (94.3%) tissues of 16 (88.9%) patients. IHC revealed a heterogeneous distribution of positive cells and stained primarily pneumocytes and alveolar macrophages and only rarely other cells such as bronchial epithelium. Quantitative analysis of SARS-CoV-2 RNA showed a significant difference between IHC positive and negative samples (101,348 viral copies per 106 RNaseP molecules vs 123, $p < 0.005$). In concordance with this finding, a wide range of viral RNA was detected in different tissue samples of each individual patient. Patients positive by IHC showed a significantly shorter duration between the diagnosis of Covid-19 to death as well as shorter hospitalization times (5.4 days vs 12.9, and 3.3 vs 13.0, $p < 0.05$, respectively).

**Discussion:** IHC can be applied to pulmonary tissue to detect SARS-CoV-2. The strong correlation between RNA load and positive IHC indicates IHC being less sensitive. However, the fact that patients with positive IHC have a significantly shorter duration of disease indicates a biological difference and supports the concept of two phases of Covid-19, an early phase driven primarily by the virus and a late phase driven by other mechanisms.
Conclusions: Our analysis indicates that scanner variability can be optimized by fine-tuning. This technique will be valuable for institutes of pathology using different scanner types.

* Student paper

P 20
Self-supervised learning for weakly labelled data in colorectal cancer*

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Background: With the recent expansion of digital pathology, the need for large scale labeled data has increased. However, data annotation is a tedious task that is both time-consuming and expensive as it involves pathologists experts. In this work, we show that we can benefit from large unlabelled cohorts to learn meaningful data representations and reduce the burden of costly annotation, closing the performance gap with supervised approaches in down-stream tasks.

Methods: We propose a self-supervised learning solution that relies on state-of-the-art momentum contrast visual data representation. The approach is based on contrastive learning that is able to discriminate between different tissue examples widely available in digital pathology. We validate our approach using two publicly available datasets of colorectal tissues that contain more than 100,000 labeled patches classified into 9 different tissue types.

Results: Firstly, we pre-train a model for each cohort without considering labels, e. i. in an unsupervised fashion. Secondly, we apply a simple linear classifier on top of frozen features (or fine-tuned) to classify the different tissue types using only a fraction of the available labels (see tables). We compare the results with the standard supervised approach. We achieve better performance on one dataset (+2.6–3.7 %) and similar performance with the second on linear evaluation. Moreover, the approach achieves a better score for specific classes such as tumors (+1.9–4.0 %), lymphocytes (+1.5–2.8 %), normal mucosa (+2.3–3.6 %), or debris (+3.2–5.4 %).

Conclusions: In our work, we showed that we could take advantage of the large quantity of unlabeled data generated by scanners to learn embedding spaces, which encode the semantic structure of data. Moreover, by labeling just a small fraction of the data, we achieve similar performances as supervised approaches. This point is critical as it would eventually reduce the annotation time for pathologists while keeping similar sensitivity-specificity trends compared to the supervised models.

* Student paper

Fig. 1 | P 20 A Kather-161 Fl-score over classes (100 % labels) for tumor (TUM), stroma (STR), complex stroma (COMP), lymphocytes (LYM), debris (DEB), normal (NORM), adipose (ADI), background (BACK)

Fig. 2 | P 20 A Kather-191 Fl-score over classes (100 % labels) for tumor (TUM), stroma (STR), mucin (MUC), lymphocytes (LYM), normal (NORM), debris (DEB), adipose (ADI), background (BACK), smooth muscle (MUS)

Fig. 3 | P 20 A Fl-score over all datasets with different percentages of training labels. We highlight top scores under statistical relevance (p-value < 0.05) with respect to other models if only part a subset of labels are used