Identification of microRNAs involved in pathways which characterize the expression subtypes of NSCLC

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Dysregulation of microRNAs is a common mechanism in the development of lung cancer, but the relationship between microRNAs and expression subtypes in non-small-cell lung cancer (NSCLC) is poorly explored. Here, we analyzed microRNA expression from 241 NSCLC samples and correlated this with the expression subtypes of adenocarcinomas (AD) and squamous cell carcinomas (SCC) to identify microRNAs specific for each subtype. Gene set variation analysis and the hallmark gene set were utilized to calculate gene set scores specific for each sample, and these were further correlated with the expression of the subtype-specific microRNAs. In ADs, we identified nine aberrantly regulated microRNAs in the terminal respiratory unit (TRU), three in the proximal inflammatory (PI), and nine in the proximal proliferative subtype (PP). In SCCs, 1, 5, 5, and 9 microRNAs were significantly dysregulated in the basal, primitive, classical, and secretory subtypes, respectively. The subtype-specific microRNAs were highly correlated to specific gene sets, and a distinct pattern of biological processes with high immune activity for the AD PI and SCC secretory subtypes, and upregulation of cell cycle-related processes in AD PP, SCC primitive, and SCC classical subtypes were found. Several in silico predicted targets within the gene sets were identified for the subtype-specific microRNAs, underpinning the findings. The results were significantly validated in the LUAD (n = 492) and LUSC (n = 380) TCGA dataset (False discovery rates-corrected P-value < 0.05). Our study provides novel insight into how expression subtypes determined with discrete biological processes may be regulated by subtype-specific microRNAs. These results may have importance for the development of combinatory therapeutic strategies for lung cancer patients.

1. Introduction

Non-small-cell lung cancer (NSCLC) accounts for approximately 85% of all lung cancers, where adenocarcinomas (AD) and squamous cell carcinomas (SCC) are the main histological subtypes (Travis, 2014). AD and SCC origin from different cell types and are associated with different types of mutations. The majority of never-smoking patients with NSCLC develop AD (Halvorsen et al., 2016; Pikor et al., 2013). The two histological subtypes can be further divided into three and four expression-based subgroups, respectively. It is

Abbreviations
AD, adenocarcinoma; EMT, epithelial–mesenchymal transition; GSE, gene set enrichment; GSVA, gene set variation analysis; PI, proximal inflammatory; PP, proximal proliferative; SCC, squamous cell carcinoma; TRU, terminal respiratory unit.
shown that the expression subtypes are robust and harbor distinct features. The ADs can be classified as terminal respiratory unit (TRU), proximal inflammatory (PI), and proximal proliferative (PP) using a nearest centroid subtype predictor of 506 genes, as previously described (Hayes et al., 2006; Wilkerson et al., 2012). The TRU subtype, including the majority of never-smokers, is associated with favorable outcome compared to patients with non-TRU AD (Ringer et al., 2016). The TRUs are recognized with low mutational burden, but with distinct driver mutations such as EGFR mutations, ALK rearrangement, and ROS1 alterations. The PI subtype is described with high immunological activity, high mutational burden, and high frequency of TP53 mutations. High frequency of TP53 mutations is also found in the PP subtype in addition to KRAS and STK11 mutations. Increased expression of DNA repair genes is reported, probably reflecting the high number of heavy smokers, found in this subgroup (Network, 2014). The SCC samples can be divided into the four expression subgroups basal, primitive, classical, and secretory based on the previously published centroid classifiers for SCC (Wilkerson et al., 2010). Tumors classified as basal are usually well differentiated and express genes involved in cell adhesion and formation of the basement membrane. The primitive subtype is associated with high proliferation, poor differentiation, and poor prognosis. The classical subtype is also recognized as an aggressive disease and is described as hypermethylated and with high chromosomal instability, probably reflecting the overall high number of heavy smokers. The secretory subtype is characterized with high immune activity and secretory functions (Network, 2012). Based on immune cell estimation, the secretory subtype shares characteristics with normal lung tissue as being immune cell-rich (Ojlert et al., 2019). Despite demonstrating distinct phenotypic and genetic differences, the molecular mechanisms underlying the development of the expression subtypes are poorly explored.

A class of small noncoding RNA called microRNA has shown to be essential in post-transcriptional regulation of mRNAs by inhibiting translation or exerting mRNA degradation (Wilczynska and Bushell, 2015). Recently, a pan-cancer project revealed the role of microRNAs in regulating gene expression signatures of the cancer hallmarks (Dhawan et al., 2018), underpinning the crucial role of microRNAs during tumorigenesis. Aberrant expression of microRNAs is well established as an important factor in lung cancer development, and dysregulation across different histological lung cancer subtypes is reported (Calin and Croce, 2006; Landi et al., 2010; Tran et al., 2018). However, the role of microRNAs in the development of the lung cancer gene expression-based subtypes is largely unknown.

In this study, we analyzed microRNA expression for a large set of NSCLC samples to identify microRNAs associated with the expression subtypes. Subtype-specific microRNA expression was correlated to gene set enrichment (GSE) scores in order to identify associated pathways that the microRNAs may be regulating, characterizing the expression subtypes. The results were further validated in independent NSCLC cohorts from The Cancer Genome Atlas (TCGA).

2. Materials and methods

2.1. Oslo cohort

Patients diagnosed with operable NSCLC from 2006 to 2014 were included in this study (n = 241). The patients underwent curatively intended surgical resection at Rikshospitalet, Oslo University Hospital, Norway. Tumor samples were snap-frozen in liquid nitrogen and stored at −80 °C until RNA isolation was performed. Clinical characteristics are outlined in Table 1. Out of the 241 samples, 132 samples were classified as ADs and 109 as SCC. Never-smokers were defined as those who had smoked < 100 cigarettes per lifetime. In this study, 19 patients diagnosed with AD were never-smokers.

The study was approved by the Regional Ethics Committee (S-05307), and written informed consent was obtained from all patients. The study was performed in agreement with the standards established by the Declaration of Helsinki.

Table 1. Patient characteristic for the Oslo cohort. Subtype refers to gene expression subtype.

| Subtype          | Adenocarcinomas (n = 132) | Squamous cell carcinomas (n = 109) |
|------------------|---------------------------|----------------------------------|
| Subtype          | n (%)                     | n (%)                           |
| PI               | 28 (21.2)                 | 30 (27.5)                       |
| PP               | 23 (17.4)                 | 6 (5.5)                         |
| TRU              | 81 (61.4)                 | 50 (45.9)                       |
| Secretory        | 23 (21.1)                 |                                 |
| Age (median)     | 66.6                      | 66.8                             |
| Gender female n | 73 (55.3)                 | 40 (36.7)                       |
| Stage n (%)      |                           |                                  |
| Ia/Ib            | 79 (59.8)                 | 56 (51.4)                       |
| Ila/Iib          | 27 (20.5)                 | 36 (33)                         |
| Ili/Iiff         | 26 (19.7)                 | 17 (15.6)                       |
| EGFR mutated n  | 19 (14.4)                 | na                               |
| Pack-years (mean)| 27.6                     | 42.4                             |
2.2. RNA extraction
Total RNA (totRNA) was extracted using standard TRIzol methods (Invitrogen, Carlsbad, CA, USA). RNA quantity and quality (yield, 260/280 ratio and 260/230 ratio) were determined using the NanoDrop ND-1000 spectrometer (NanoDrop Technologies, Wilmington, DE, USA). The RNA integrity numbers (RIN) were assessed using Agilent 2100 Bioanalyzer and totRNA Nano Kit (Agilent Technologies, Santa Clara, CA, USA) according to the manufacturer’s protocol.

2.3. mRNA expression analyses
We analyzed mRNA expression from the tumor samples using gene expression microarray from Agilent Technologies (SurePrint G3 Human GE, 8 × 60 K). For the AD samples, we used v.1, whereas for the SCC samples, we used v.3 of the microarray platform. We used 50 ng totRNA as input for the analyses, and the analyses were performed according to the protocol from the supplier. The data are deposited at ArrayExpress with accession number: E-MTAB-7954.

2.4. microRNA expression analyses
We analyzed microRNA expression using Agilent Human microRNA Microarray for 132 ADs (microarray kit release 16.0, 8 × 60 K) and 109 SCCs (microarray kit release 21.0, 8 × 60 K). We used 100 ng of totRNA in the analyses following the protocol as specified by the manufacturer. The data are deposited at ArrayExpress with accession number: E-MTAB-7958.

2.5. Normalization of data
Data from the microRNA analyses were log2-transformed and normalized using the 90th percentile method. The gene expression data were log 2-transformed and quantile normalized in GENESPRING GX Analysis Software v.12.1 (Agilent Technologies). We filtered out microRNAs detected in <10% of the AD samples and in <20% of the SCC samples. After filtering, 562 and 905 microRNAs remained for further analysis, respectively.

2.6. Molecular subtyping of adenocarcinomas and squamous cell carcinomas
The AD samples were assigned a gene expression subtype being TRU, PP, or PI, using the previously described 506 gene centroid classifier and Pearson correlation (Wilkerson et al., 2012). The SCC were classified as basal, secretary, primitive, or classical based on the centroid classifier described for SCCs (Wilkerson et al., 2010). Samples negatively correlated with all subtypes were not assigned to any subtype.

2.7. Validation dataset
For validation, the lung AD (LUAD) and the lung SCC (LUSC) datasets were obtained from TCGA (Network, 2012, 2014). microRNA and mRNA expression data were extracted as log2(RPKM + 1) values through the Xena browser (https://xenabrowser.net/datapages/). Expression subtyping and gene set variation analysis (GSVA) were performed on mRNA sequencing data from the LUSC (n = 553) and the LUAD dataset (n = 576). Results from the microRNA analysis were validated in 492 LUAD samples and 380 LUSC samples extracted from TCGA. In addition, we included 45 normal lung tissue samples from the LUAD dataset and 44 normal tissue samples from the LUSC dataset.

2.8. Statistics
All statistics were done in R version 3.5.2 (R Development Core Team, 2013). Hierarchical clustering was performed with ComplexHeatmap package version 1.20.0 using ward.D2 as clustering method (Gu et al., 2016). Kruskal–Wallis tests were applied to identify microRNAs differentially expressed between the expression subtypes. Following a significant Kruskal–Wallis test, a post hoc Dunn test was utilized to pinpoint in which subtype the microRNA was differentially expressed compared to the others. The packages FSA, fisheries stock analysis R package version 0.8.22 FSA v0.8.22, and Reshape (Wickham, 2007) (Ogle et al., 2018) were utilized. False discovery rates (FDR) were controlled using Benjamini–Hochberg adjustment (Yoav Benjamini, 1995). FDR-corrected P-values < 0.05 were assigned statistically significant.

Gene set variation analysis is a nonparametric and unsupervised method for assessing GSE in gene expression data. This method allows the evaluation of pathway enrichment for each sample (Hanzelmann et al., 2013). Here, we used the R package GSVA with the Molecular Signatures Database (MSigDB) hallmark gene sets (n = 50) downloaded from Broad Institute (http://software.broadinstitute.org/gsea/msigdb/collections.jsp#H) to assess enrichment in the samples. The hallmark gene set contains specific well-defined biological states or processes and displays coherent expression (Liberzon et al., 2015). An enrichment score for each of the 50 processes was calculated for each sample. Then, Spearman rank correlation was assessed.
between the enrichment scores and each of the differentially expressed microRNAs. Bonferroni adjustment was applied to the correlation values to correct for multiple testing. Corrected \( P \)-values < 0.05 were assigned as statistically significant and further considered. We used miRDIP 4.1 (http://ophid.utoronto.ca/miRDIP/) to identify predicted targets for the subtype-specific microRNAs (Tokar et al., 2018). This database integrates several computational microRNA–target prediction tools aiming to strengthen the prediction of microRNA/target relationship. Only predictions ranked as very high, corresponding to the top 1% of the list, were accepted as potential targets. Gene sets positively correlated with subtype-specific microRNAs were not tested. Further, only gene sets with the highest anticorrelated pathway for each microRNA in both cohorts were selected for prediction analyses. Genes were identified as targets for the tested microRNAs if the correlation coefficients were significantly negative (Bonferroni-corrected \( P \)-value < 0.05) in both cohorts.

3. Results

The frequencies of the AD subtypes TRU, PI, and PP detected in the Oslo cohort were in line with the TCGA LUAD cohort, although more TRU samples (61.4% versus 45.8%) and fewer PI samples (21.2% versus 35.3%) were identified (Fig. 1). For the SCC samples, a lower frequency of the primitive subtype was detected in the Oslo cohort (5%) than in the TCGA LUSC cohort (14%) as shown in Fig. 1.

We identified 251 microRNAs differentially expressed between the expression subtypes in ADs in the Oslo cohort (FDR-corrected Kruskal–Wallis \( P \)-value < 0.05). Of these, 157 microRNAs were validated in the LUAD cohort of TCGA (FDR-corrected Kruskal–Wallis \( P \)-value < 0.05). Dunn’s test was subsequently used to find expression subtype-specific microRNAs. In order to be classified as subtype-specific, the level had to be significantly different from the other subtypes (FDR-corrected \( P \)-value < 0.05) in both cohorts (Oslo and TCGA). As shown in Fig. 2, the number of microRNAs expressed at different levels was highest (low \( P \)-values are displayed in blue) when we compared PP and TRU samples. The PI and PP samples showed a more similar microRNA expression pattern.

Next, we included normal lung tissue samples (LUAD, \( n = 45 \)) and focused only on subtype-specific microRNAs that were also differentially expressed compared to the normal samples. Most of the microRNAs that had similar level as the normal samples were associated with the TRU subtype. These were filtered out. Following these criteria, 21 subtype-specific microRNAs were identified in ADs (Table 2, Fig. S1, Table S1): three microRNAs characterizing PI (all up), nine microRNAs characterizing PP (two up and seven down), and nine microRNAs characterizing TRU (all up).

We applied the same analysis to the SCC samples. Using Kruskal–Wallis test, we identified 50 microRNAs being differentially expressed between the expression subtypes in the Oslo cohort (FDR-corrected \( P \)-value < 0.05). Of these, 41 were validated in the LUSC cohort (FDR-corrected \( P \)-value < 0.05). Dunn’s tests further excluded 21 microRNAs, leaving 20 microRNAs passing the above-mentioned criteria (Table 2, Fig. S1, Table S1): one microRNA characterizing basal (up), five microRNAs characterizing classic (all up), five microRNAs characterizing primitive (three up and two down), and nine microRNAs characterizing secretory (one up and eight down). Of note, due to the low number of primitive samples in the Oslo cohort, borderline significant microRNAs for tests with this subtype were included if significant in the LUSC cohort. As shown in Fig. 2, the largest difference between the SCC subtypes, in terms of microRNA expression, was found between the secretory and the classical subtypes. On the other side, the classical and the primitive subtypes were most similar with regard to microRNA expression (Fig. 2).

3.1. Gene set variation analysis

Gene set variation analysis is a GSE method that estimates underlying pathway activity variation in samples in an unsupervised manner. The hallmark gene sets from MSigDB were used for the analysis (Liberzon et al., 2015). These gene sets contain 50 well-defined signatures of 50 hallmarks that represent well-defined biological processes. An enrichment score was calculated sample-wise for each hallmark without knowledge of any phenotypic information. In order to identify hallmarks associated with subtype-specific microRNAs, the enrichment scores were correlated with the expression level of the subtype-specific microRNA using Spearman rank correlation (retaining correlation with a Bonferroni-corrected \( P \)-value < 0.05). As shown in Table 3, immune response, cell cycle maintenance, epithelial–mesenchymal transition (EMT), and metabolism were the processes that correlated the most with the subtype-specific microRNAs. More details are shown in Table S2.

To further explore and visualize the correlation between subtype-specific microRNAs and hallmark signatures, the correlation values were hierarchically clustered (Figs 3 and S2).
As displayed in Fig. 3 and Table 3, the AD PI subtype shows upregulation of processes involved in immune response and of the hallmarks cell cycle and DNA repair, which were the opposite of what was found in AD PP subtype. The TRU subtype was associated with upregulation of bile acid metabolism and downregulation of cell cycle and DNA repair. For SCC, a similar pattern was seen with immune response upregulated and cell cycle and DNA repair downregulated in the secretory subtype, just the opposite of what was detected for the primitive and classical subtypes.

In order to assess whether the subtype-specific microRNA signal originates from the lung cancer cells or from infiltrating immune cells, we compared our subtype-specific microRNAs with the results from a study investigating human cell-specific microRNA expression. In this project, the authors sequenced microRNAs from 46 primary cell types, 42 cancer cell lines and tissues (McCall et al., 2017). We extracted the sequencing data from dendritic cells, B lymphocytes, T lymphocytes, macrophages, blood, lung fibroblasts, lung tissue, and lung cancer cell lines. All over, most of the subtype-specific microRNAs showed highest expression in lung tissue and lung cancer cell lines. Nevertheless, miR-142-3p, miR-142-5p, and miR 146-5p were highly expressed in T cells, whereas miR-140-3p and miR-221-5p were highly expressed in B cells. We also found microRNAs which seemed to be exclusively expressed in lung tissue and lung cancer cell lines. This included 149-5p, miR-196a-5p, miR-200b-3p, miR-224-5p, miR-429, and miR-452-5p. (Fig. S3, Table S3).

3.2. Prediction of targets for the subtype-specific microRNAs

In order to further elucidate how the subtype-specific microRNAs may regulate the associated gene sets, we utilized prediction analysis to find potential targets for the microRNAs within the gene sets. First, we identified gene sets being anticorrelated with the subtype-specific microRNAs. The most anticorrelated gene set for each of the subtype-specific microRNAs being significant in both cohorts was selected for target analysis. One exception was made; we included Spermatogenesis and E2F targets for the TRU subtype since these two gene sets were anticorrelated with several of the TRU-specific microRNAs, but were not ranged as the most anticorrelated gene sets. As expected, for the AD PI, AD TRU, SCC basal, and SCC classical subtypes, no upregulated gene sets were anticorrelated with the selected microRNAs due to only upregulated microRNAs. For AD PP, 11 predicted targets were identified for miR-101-3p and miR-140-3p within the gene set G2M checkpoint, and six predicted targets within inflammatory response were significantly anticorrelated with miR-200c-3p.
and miR-141-3p. For SCC secretory subtype, predicted targets for miR-200a-3p, miR-200b-3p, miR-200c-3p, miR-141-3p, miR-205-5p, miR-429, and miR-196-5p were identified within the gene sets inflammatory response (12 predicted targets), EMT (32 predicted targets), myogenesis (16 predicted targets), kras signaling up (13 predicted targets), il6-jak-stat3-signaling (four predicted targets), and/or coagulation (12 predicted targets). The upregulated gene sets in the SCC primitive subtype E2F targets and G2M checkpoint were identified with eight and 11 predicted targets for miR-22-3p and miR-145-5p, respectively. Further, four predicted targets for miR-106-5p were found in the downregulated gene sets myogenesises and coagulation. Three downregulated gene sets were identified with predicted targets for the SCC classical subtype. For more details, see Table S4. Nine of our predicted targets have been functionally validated in previous studies, according to MiRTargetBase (Chou et al., 2018). Details are shown in Table S4.

4. Discussion

In this study, we identified 21 and 20 subtype-specific microRNAs in AD and SCC, respectively. Correlation analysis between microRNA expression and hallmark enrichment scores revealed distinct positive and negative associations with the subtypes of AD and SCC. This suggests that the identified microRNAs may regulate biological processes determining the different subtypes. Even though the identified microRNAs were subtype-specific, they were involved in many of the same processes, although associated with different targets. We propose that distinct processes characterizing the subtypes, such as high immune activity in AD PI and SCC secretory samples, and proliferation in AD PP, SCC primitive, and SCC classical samples, may be
regulated by the identified subtype-specific micro-RNAs.

4.1. The role of microRNAs in cell cycle and proliferation

Pathways involved in cell cycle and DNA repair signaling were upregulated in SCC classical, SCC primitive, and AD PP subtypes and downregulated in AD TRU and SCC secretory subtypes. These results are in line with previous work where a high proliferation score was detected in the classical, primitive, and PP subtypes (Ojlert et al., 2019). In the SCC classical subtype, all the five classical-specific microRNAs were upregulated, and subsequently, no anticorrelated predicted targets were detected within the upregulated gene sets. Therefore, we speculate that targets for these microRNAs are inhibitors of the G2M checkpoint and E2F target gene sets, leading to an increased signaling. For the primitive tumors, the same pathways were upregulated but seem to be controlled in a different manner. With 19 predicted targets of miR-22-3p and miR-145-5p within G2M checkpoint and E2F target gene sets, an upregulation of these gene sets may be explained by downregulation of the microRNAs controlling these processes.

In the AD PP subtype, we found 11 predicted targets of miR-140-3p and miR-101-3p within the G2M checkpoint gene set indicating that upregulation of G2M checkpoint most likely is a result of repression of miR-140-3p and miR-101-3p. However, since miR-101-3p is expressed both in immune cells and lung tissue, they may be targets of essential components within the cell cycle, simultaneously being involved in the development of suppressive mechanisms in the immune microenvironment. Furthermore, since miR-140-3p is highly expressed by B cells, low levels of miR-140-3p may indicate the absence of B-cell infiltration. The TRU tumors were associated with a downregulation of G2M checkpoint and E2F-targets. High levels of miR-181c-5p may explain this downregulation, supported by identification of 12 predicted targets within these two signaling pathways. None of the 12 predicted targets have been functionally validated, but a recent study showed that miR-181c-5p is involved in G2M-checkpoint regulation, and have direct targets within this pathway (Sun et al., 2019).

We found an upregulation of the DNA repair in primitive, classical, and PP subtypes. In addition, the gene set reactive oxygen species were upregulated in the classical subtype. This probably reflects that these subtypes are associated with more heavy smoking (Wilkinson et al., 2010; Wilkerson et al., 2012).
### Table 3. Hallmark gene sets significantly associated with high or low expression of subtype-specific microRNAs in both the Oslo and TCGA cohorts.

| Upreg | AD | PI | AD | PP | AD | TRU |
|-------|----|----|----|----|----|-----|
| allograft_rejection | dna_repair | bile_acid_metabolism |
| complement | g2m_checkpoint | |
| il2_stat5_signaling | myc_targets_v1 | |
| inflammatory_response | myc_targets_v2 | |
| interferon_alpha_response | glycolysis | |
| interferon_gamma_response | oxidative_phosphorylation | |
| tnfa_signaling_via_nfk | | |

| Downreg | AD | PI | AD | PP | AD | TRU |
|---------|----|----|----|----|----|-----|
| allograft_rejection | e2f_targets | |
| apoptosis | g2m_checkpoint | |
| hedgehog_signaling | spermatogenesis | |
| il2_stat5_signaling | myc_targets_v1 | |
| inflammatory_response | myc_targets_v2 | |
| interferon_alpha_response | unfolded_protein_response | |
| interferon_gamma_response | mitotic_spindle | |
| tnfa_signaling_via_nfk | mtorc1_signaling | |
| p53_pathway | glycolysis | |

| Upreg | SCC Primitive | SCC Classical | SCC Secretory | SCC Basal |
|-------|---------------|---------------|---------------|-----------|
| dna_repair | dna_repair | allograft_rejection | glycolysis | estrogen_response_late |
| e2f_targets | p53_pathway | inflammatory_response | | |
| g2m_checkpoint | reactive_oxygen_species_pathway | il6_jak_stat3_signaling | | |
| prog_cell_cycle | e2f_targets | complement | | |
| mitotic_spindle | g2m_checkpoint | epithelial_mesenchymal_transition | | |
| spermatogenesis | unfolded_protein_response | angiogenesis | | |
| myc_targets_v1 | spermatogenesis | coagulation | | |
| myc_targets_v2 | myc_targets_v1 | myogenesis | | |
| oxidative_phosphorylation | myc_targets_v2 | kras_signaling_up | | |
| fatty_acid_metabolism | | | | |
| glycolysis | | | | |
| oxidative_phosphorylation | | | | |
| cholesterol_homeostasis | | | | |
| peroxisome | | | | |
| mtorc1_signaling | | | | |
| estrogen_response_late | | | | |

| Downreg | SCC Primitive | SCC Classical | SCC Secretory | SCC Basal |
|---------|---------------|---------------|---------------|-----------|
| allograft_rejection | allograft_rejection | dna_repair | NA | |
| complement | inflammatory_response | g2m_checkpoint | |
| inflammatory_response | complement | e2f_targets | |
| il2_stat5_signaling | interferon_alpha_response | unfolded_protein_response | |
| inflammatory_response | interferon_gamma_response | spermatogenesis | |
| il6_jak_stat3_signaling | myc_targets_v1 | myc_targets_v2 | |
| epithelial_mesenchymal_transition | myc_targets_v2 | fatty_acid_metabolism | |
| angiogenesis | mtorc1_signaling | cholesterol_homeostasis | |
| coagulation | kras_signaling_up | metabolic_pathway | | |
| heme_metabolism | | | | |
| apoptosis | | | | |
4.2. Immune activity regulated by microRNAs

Processes belonging to immunological response were upregulated in AD PI and SCC secretory subtypes and downregulated in AD PP, SCC primitive, and SCC classical subtypes. Interestingly, all three microRNAs associated with the PI subtype are highly expressed in B cells and T cells (Fig. S3), which point to a high

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**Table 3.** (Continued).

| Downreg | SCC Primitive | SCC Classical | SCC Secretory | SCC Basal |
|---------|---------------|---------------|---------------|-----------|
| uv_response_dn | myogenesis | kras_signaling_up | apical_junction | androgen_response |

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**Fig. 3.** shows the correlation between the subtype-specific microRNAs and the hallmark gene set for AD and SCC in the Oslo cohort. Subtype annotation indicates which subtype the different microRNAs are associated with. To identify up- or downregulated pathways, the correlation coefficient for downregulated microRNAs (annotated with black/low) must be multiplied with −1 (this will switch the red pixels into blue and vice versa).

4.2. Immune activity regulated by microRNAs

Processes belonging to immunological response were upregulated in AD PI and SCC secretory subtypes and downregulated in AD PP, SCC primitive, and SCC classical subtypes. Interestingly, all three microRNAs associated with the PI subtype are highly expressed in B cells and T cells (Fig. S3), which point to a high
immune activity in the PI subtype. This is in line with previous work where high cytolytic and immune score in the PI and secretory subtypes, and low cytolytic and immune score in the PP, primitive, and classical subtypes were detected (Ojlert et al., 2019). Interestingly, in a study of microdissected lung ADs, genes involved in the immune response were not as prominent in the PI subtype as we detected in our study. One possible explanation is that we used bulk tumor, which also harbors cells from the microenvironment such as immune cells (Zabeck et al., 2018). This indicates that the expression subtypes are not restricted to the tumor cell, but can also mirror the tumor microenvironment.

The secretory subtype was recognized with upregulation of immune-related pathways, which is confirmed in other studies (Faruki et al., 2017; Ojlert et al., 2019). However, in contrast to the PI subtype, the subtype-specific microRNAs detected in secretory tumors were downregulated (except for miR-30c-2-3p) and mostly associated with expression in lung tissue. Since the secretory subtype is reported to be associated with high immune activity, our findings may indicate that these microRNAs are expressed by the tumor cells, usually suppressing the immune activity. This was further supported when we identified 16 predicted targets within the gene sets Immune response and il6-jak-stat3-signaling. Interestingly, this was also consistent with the finding of a downregulation of immune response in the AD PP subtype, identified with upregulation of two of the same microRNAs (miR-200c-3p and miR-141-3p) which were downregulated in the SCC secretory subtype.

The expression pattern for the secretory subtype shares many similar features to the expression pattern of normal lung tissue. Both normal lung tissue and secretory lung cancer tissue exhibit an immune active pattern. This may explain why many of the secretory subtype-specific microRNAs were in the same direction as the normal samples.

4.3. Epithelial–mesenchymal transition signaling

Several studies have shown that members of the miR-200 family (miR-200a, miR-200b, miR-200c, miR-429, and miR-141) are crucial regulators of the EMT signaling (Humphries and Yang, 2015). It has been shown that miR-200 family can target ZEB1 and ZEB2 and promote expression of E-cadherin, thus hindering migration, invasion, and tumor angiogenesis (Korpel et al., 2008). Tumors with the secretory subtype were recognized with a downregulation of all members of the miR-200 family and upregulation of EMT and angiogenesis. For EMT signaling, 32 predicted targets for the miR-200 family were identified within our dataset supporting these findings. Interestingly, in a study of pan-cancer EMT signature, immune cell signaling was strongly correlated to EMT (Mak et al., 2016). This was also shown with the clustering of microRNAs and the hallmark gene sets (Fig. 3) where EMT- and immune-related processes were identified in the same subcluster. This may explain why the classical and primitive subtypes revealed a downregulation of EMT and angiogenesis, in addition to a low immune activity. An opposite result was discovered for the SCC secretory subtype, identified with upregulation of the same pathways. Further, an association between EMT and increased expression of PD-L1 has been reported, and EMT has been suggested in regulating immune escape in lung cancer (Chen et al., 2014). However, we did not observe high PD-L1 expression in the secretory subtype in our previous study (Ojlert et al., 2019).

There are some limitations in this study. Due to fewer samples included in the Oslo cohort, microRNAs with borderline significance were included if significant in the TCGA cohort. This may imply that other microRNAs important for this subtype were not captured during the first analysis. However, all the results were validated in TCGA which make the findings in this study robust and repeatable. Furthermore, only one microRNA (miR-31-5p) was significantly associated with the basal subtype, resulting in few pathways correlated with this group. This may indicate that development of basal tumors is driven by other mechanisms than microRNAs, or that some basal-specific microRNAs were not captured by our analyses. However, we found that miR-31-5p was highly lung cancer and lung tissue specific which may indicate that this microRNA is oncogenic and plays a specific role in basal lung tumors.

The samples from the Oslo cohort were analyzed using a microarray platform, whereas the TCGA samples were profiled using next-generation sequencing technology. Thus, there may exist additional subtype-specific microRNAs which were not captured in this study as the microRNAs present on the microarray defined the microRNA focus. Nevertheless, the resulting subtype-specific microRNAs reported in the present study were robustly identified across platforms.

5. Conclusions

In this study, we showed that subtype-specific microRNAs may be involved in essential processes characterizing the expression subtypes of ADs and SCCs. Of
note, functional studies are warranted in order to detect precise targets for the subtype-specific microRNAs. Unraveling the underlying biology in lung cancer subtypes may be important in order to offer the patients a more stratified targeted therapy. Inhibition of essential pathways together with standard care of treatment including immunotherapy may be a beneficial strategy.

Acknowledgements

The authors want to thank Ingjerd Solvoll for help with collecting tissue samples and clinical information. The authors gratefully acknowledge the contribution from the patients participating in this study. This work was supported by The Norwegian Cancer Society (grant number 88503-2013) and South-Eastern Norway Regional Health Authority (grant number 2016056).

Conflict of interest

The authors declare no conflict of interest.

Author contributions

MRA, AH, and ARH conceived and design the project. ARH acquired the data. ARH, MRA, DN, and AKØ analyzed and interpreted the data. ARH, AH, and MRA wrote the paper. SS, AH, and OTB collected the clinical data and the material, and participated in the evaluation of the data. All authors read, revised the manuscript critically, and approved the final manuscript.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1.** Examples of microRNAs significantly associated with one specific expression subtype shown for the LUAD and LUSC cohorts.

**Fig. S2.** This figure shows the correlation between the subtype-specific microRNAs and the hallmark gene set for AD and SCC in the TCGA cohort. Subtype annotation indicates which subtype the different microRNAs are associated with. To identify up- or down-regulated pathways, the correlation-coefficient for downregulated microRNAs (annotated with black/low) must be multiplied with -1 (this will switch the red pixels into blue and vice versa).

**Fig. S3.** This figure shows the expression of the identified subtype-specific microRNAs across different immune cells, lung tissue and lung cancer cell lines. Data is extracted from McCall et al. The color-bars show the subtype and direction associated with the microRNA. Three microRNAs (miR-141-3p, miR-145-5p, miR-200c-3p) were specific to subtypes both in AD and SCC, and are annotated with the suffix 2.

**Table S1.** Dunn test was utilized to identify expression subtype-specific microRNAs in Oslo Cohort and TCGA.

**Table S2.** Shows the hallmark gene sets associated with subtype-specific microRNAs.

**Table S3.** Shows which cells the subtype-specific microRNA may originate from. The sequencing data is from a project were the authors sequenced microRNAs from 46 primary cell types, 42 cancer cell lines and tissues (McCall et al., 2017). We extracted the sequencing data for the subtype-specific microRNAs.

**Table S4.** Shows in silico predicted targets for subtype-specific microRNAs within the associated gene set. Genes marked with a star are previous functionally validated according to MiRTargetBase (Chou et al., 2018).