The transcriptome-wide landscape of molecular subtype-specific mRNA expression profiles in acute myeloid leukemia

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Abstract
Molecular classification of acute myeloid leukemia (AML) aids prognostic stratification and clinical management. Our aim in this study is to identify transcriptome-wide mRNAs that are specific to each of the molecular subtypes of AML. We analyzed RNA-sequencing data of 955 AML samples from three cohorts, including the BeatAML project, the Cancer Genome Atlas, and a cohort of Swedish patients to provide a comprehensive transcriptome-wide view of subtype-specific mRNA expression. We identified 729 subtype-specific mRNAs, discovered in the BeatAML project and validated in the other two cohorts. Using unique proteomics data, we also validated the presence of subtype-specific mRNAs at the protein level, yielding a rich collection of potential protein-based biomarkers for the AML community. To enable the exploration of subtype-specific mRNA expression by the broader scientific community, we provide an interactive resource to the public.

1 | INTRODUCTION

Acute myeloid leukemia (AML) is a heterogeneous disease due to the diversity of genetic alterations.1,2 These alterations determine AML pathophysiology, progression and response to therapy, and thus disease heterogeneity complicates clinical management.3-4 Many studies have aimed to stratify AML patients into molecular subtypes, each with distinct clinical significance in terms of prognosis or therapy response.5-7 For example, favorable outcomes are reported for patients with fusion subtypes, such as PML-RARA and CBFB-MYH11, as well as the non-fusion subtypes, such as CEBPA biallelic subtype.6,8 On the other hand, overexpression of EVI1 is a poor prognostic factor in MLL-rearranged AML.9 A fully genomic classification of AML that considers patterns of co-mutations has been established recently...
based on a large patient cohort. At least 11 molecular subtypes can be recognized based on cytogenetic and targeted sequencing analyses. However, large-scale information on how these genetic differences manifest themselves at the transcriptome and proteome level is not yet established. Therefore, our aim is to identify transcriptome-wide mRNAs that are specific to each of the molecular subtypes, and to validate their significance at the protein level. Subtype-specific mRNAs or the corresponding protein can provide further biological understanding and be utilized as potential biomarkers for diagnosis, classification, therapy response, as well as disease monitoring.

Here, we use a novel method that leverages available large-scale RNA-seq datasets to identify molecular subtype-specific mRNA expression patterns. The specificity of the genes is established using two statistics, one for testing the genes that are over-expressed in a single subtype compared to all the other subtypes and the other for testing whether the remaining subtypes are statistically different.

We analyzed next-generation RNA-sequencing (RNA-seq) data of 955 AML samples from three cohorts, including the BeatAML project (N = 461), the Cancer Genomic Atlas (N = 179) and a cohort of Swedish patients (N = 315), called the Clinseq cohort from here on. To provide these results and findings to the broader scientific community, we have created an interactive resource containing the complete results of subtype-specific analysis at https://ngiavtr.shinyapps.io/AMLSubtypeSpecificDiscovery/.

2 | MATERIALS AND METHODS

2.1 | Study cohorts

The statistical identification of subtype-specific genes is optimized using several tuning parameters, including the choice of contrast statistics and their corresponding thresholds. The BeatAML cohort which contains 461 AML patients is used as a discovery cohort to determine subtype-specific genes. The TCGA cohort (179 samples), and the Clinseq cohort (315 samples) are used for validation. The details of each cohort are described below. The FASTQ files were obtained from the three cohorts and RNA-seq reads were aligned to the human genome hg19. Also, XAEM was used to obtain the gene expression (transcript per million - TPM) from the RNA-seq data. The calculation process of XAEM followed the instructions provided at http://fafner.meb.ki.se/biostatwiki/xaem/.

2.1.1 | BeatAML data

The BeatAML project generated functional genomic data of primary bone marrow biopsies from patients with AML. The dataset includes genomic and transcriptomic analyses, clinical annotations and drug responses. Patients in this cohort have received standard intensive chemotherapy analogous to TCGA and Clinseq cohorts. Samples were first processed with the Agilent SureSelect Strand-Specific RNA Library Preparation Kit on the Bravo robot (Agilent). Sequencing was performed on the Illumina HiSeq 2500 platform using a 100-cycle paired-end protocol. More details of the data can be found in the original paper. In total 23 360 genes across 461 samples with complete clinical information and drug response are used in this study.

2.1.2 | TCGA data

The systematic study of the Cancer Genome Atlas (TCGA) AML samples has provided a genomic landscape of AML and generated a catalogue of leukemia-related genes. There is, therefore, a possibility to also make use of this sequencing data for a more refined understanding of subtype-specific patterns of mRNA expression. RNA-seq data was obtained using Illumina HiSeq 2000 PE 75 base sequencing protocol. Patients of the TCGA-AML study received intensive induction treatment (chemotherapy). A total of 22 374 genes from the 179 samples were used in this study as a validation set.

2.1.3 | Clinseq data

The Clinseq cohort includes 315 patients diagnosed with AML in Sweden between February 1997 and August 2014. Clinical information was retrieved from patient records and the Swedish Adult Acute Leukemia Registry. All patients underwent intensive induction therapy (including anthracyclines and cytosine arabinoside) as first-line treatment. Bone marrow or peripheral blood samples were obtained at the time of diagnosis, separated for mononuclear cells and stored at −180°C until use. Transcriptomic RNA was sequenced using the Illumina HiSeq-2500 platform. As a validation set, gene expressions of 23 572 genes across 315 samples are considered in the analysis.

A representative subset of 118 patient samples with sufficient biological material was selected from the original cohort for mass spectrometry-based proteomics analysis. The corresponding protein levels of 12 142 genes were quantified in at least one sample. Mass spectrometry based proteomics was carried out as previously described. Briefly, viably frozen patient samples were washed, and the cells were lysed by 4% SDS lysis buffer and prepared for mass spectrometry analysis using a modified version of the SP3 protein clean up and digestion protocol. Peptides were labeled with TMT10-plex reagent according to the manufacturer’s protocol (Thermo Fisher Scientific) and separated by immobilized pH gradient−isoelectric focusing (IPG-IEF) on 3–10 strips as described previously. Extracted peptide fractions from the IEF-IEF were separated using an online 3000 RSLCnano system coupled to a Thermo Fisher Scientific Q Exactive-HF. MSGF+ and Percolator in the Nextflow platform were used to match MS spectra to the Ensembl92 human protein database.
Papamannouil's classification compartmentalized AML into 11 subtypes, each with distinct diagnostic features and considerable relevance to clinical outcomes. Although the 11 subtypes well reflect the genetic characteristics of AML, the association between molecular mutations and the gene expression level are not well described.

We applied this genomic classification to the BeatAML, TCGA and Clinseq cohorts and found similar frequencies of membership in each subtype, shown in Table S1. For simplicity, in this context, we named each subtype as follows:

- NPM1—AML with NPM1 mutation;
- TP53-mutant—AML with TP53 mutation, chromosomal aneuploidy or both;
- Splice—AML with mutated chromatin, RNA-splicing genes, or both;
- CBFB-MYH11—AML with inv(16)(p13.1q22) or t(16;16)(p13.1;q22);
- PML-RARA—AML with t(15;17)(q22;q12);
- MLL—AML with MLL fusion genes, t(x;11)(x;q23) multiple fusion partners for MLL;
- CEBPA<sup>Bi</sup>—AML with biallelic CEBPA mutations;
- RUNX1-RUNX1T1—AML with t(8;21)(q22;q22);
- IDH2<sup>r172</sup>—AML with IDH2<sup>r172</sup> mutations and other class-defining lesions;
- Inv(3) with inv(3)(q21q26.2) or (3;3)(q21;q26.2);
- Other—samples that are not assigned to any of above subgroups.

More detailed information of this genomic classification for the multiple mutations and “other” group are described in the supplementary document.

### 2.3 Systematic identification of subtype-specific genes

A subtype-specific gene must be over-expressed in a single subtype compared to all the other subtypes while the other subtypes are not statistically different from each other. However, the problem is complicated due to the fact that we have 11 subtypes. If we just test the difference between one subtype against the rest, it only implies that this subtype is different from the combined distribution of the rest but the other subtypes could also different from each other, such that the specificity to one single subtype cannot be guaranteed.

To overcome this issue, we apply a method that is originally described in<sup>19</sup> and use 11 AML subtypes, as described in section 2.2, for identifying subtype-specific genes. This method computes two statistics— a robust t test (T1) and chi-square test (T2) for each subtype. So, T1 is used to determine if there is a significant difference between each single subtype and all other subtypes. And, T2 is used to test if the other subtypes have similar expression. To be considered as subtype-specific, the statistic of T1 must be large so that the subtype is significantly different from the others, but the statistic of T2 must be small to control the similarity between the remaining subtypes. The statistical significance is expressed in terms of false discovery rate (FDR) to account for multiple testing.<sup>20</sup> Following,<sup>19</sup> we set the threshold of T1-based FDR < 0.01 and T2-based FDR > 0.10.

### 2.4 Code availability

The codes to compute two statistics T1 and T2 for identifying subtype-specific genes used in this study are adapted from the original study<sup>19</sup> and can be downloaded from the public Zenodo repository at https://doi.org/10.5281/zenodo.4036552.

### 3 RESULTS

The pipeline of the subtype-specific genes discovery analysis is illustrated in Figure 1. Further experimental details are described in materials and methods section. Table S1 shows the number of patients in each subtype; NPM1 is the largest group across three cohorts (including 22.5% of samples), which is in agreement with previous reports.<sup>6,21</sup> The TP53-mutant subtype is of similar proportion to the NPM1 subtype, and the splice subtype is in the 3rd largest group. These three subtypes contain more than half of samples, accounting for 54.8% of the three cohorts. Some small subgroups, that is, CBFB-MYH11, PML-RARA, RUNX1-RUNX1T1, Inv(3) and MLL fusions, represent less than 5% of the three cohorts, in line with the WHO classification.

The complete result for all genes can be found at an interactive website https://nghiavtr.shinyapps.io/AMLSubtypeSpecificDiscovery/.

#### 3.1 Using this resource to enable the discovery of subtype-specific genes

As an example of gene-level distributions, Figure 2 presents the boxplots of the gene-level mRNA expression of PTPRG, a top-ranking gene specific to the PML-RARA subtype. It shows that PTPRG expression patterns are similar across the three cohorts, with high expression in the PML-RARA and low expression in the other subtypes, which is in agreement with other analysis.<sup>22,23</sup> The statistics of our method to identify subtype-specific genes are: T1 ≈ 0 (t-statistic = 73.56), T2 ≈ 1 (Table S2). The PTPRG expression is also validated at the protein level with higher protein levels in PML-RARA compared to all the other subtypes, as shown in Figure 2D. The top five genes in each subtype are given in Table S2. To select these, we first filter out the genes with T2-based FDR < 0.1, then rank the genes by T1.

Figure S1 shows the number of subtype-specific genes assigned to each subtype based on the BeatAML as the discovery set. There were 9226 subtype-specific genes across the 11 subtypes, with the NPM1 subtype being the largest group with 3687 genes (40%). We also investigate if the genes specific to the NPM1 subtype are able to separate patients with and without FLT3-ITD (+/−) within this subtype. Figure S2 shows a tSNE plot of mRNA expressions of the top 25 subtype-specific genes for the two groups of FLT3-ITD +/−. The result shows no separation between two groups, indicating these subtype-specific genes are specific to NPM1 but do not contain signals for the FLT3-ITD status. The subtype “other” is the smallest group including only 52 genes (0.6%).
which is reasonable since this group has no particular criteria associated. The distribution of T1 of the identified subtype-specific genes for each subtype based on the BeatAML dataset are shown in Figure S3.

3.2 Validation of subtype-specific genes

Figure S4 shows the Venn diagrams the subtype-specific genes from the three cohorts for each subtype. In total, 729 subtype-specific genes discovered in the BeatAML cohort are validated in both the TCGA and Clinseq cohorts. The NPM1 and PML-RARA subtypes have significantly higher numbers of overlapping genes across the three cohorts, with 240 and 210 respectively, compared to other subtypes (≤110).

In the heat map (Figure 3A) displaying the top 25 genes from each subtype in the discovery set (BeatAML), a pattern that distinguishes the different subtypes is evident. For patients with the NPM1, CEBPA Biallelic, CBFB-MYH11, PML-RARA, MLL and splice subtypes, these data suggest gene expression patterns specific to the corresponding subtypes. Similar patterns are observed in the validation sets (Clinseq and TCGA) for the NPM1, CEBPA Biallelic, CBFB-MYH11 and PML-RARA subtypes, as shown in Figure 3B, C. Taking the NPM1 subtype as an example, the top 25 subtype-specific genes are clearly more highly expressed in the patients classified in the NPM1 subtype than in the other subtypes. Furthermore, this pattern is also observed in the validation sets. Since the expression data of some genes are not provided in the TCGA data set, we indicate these absent genes with white lines (Figure 3C). Based on the clinical information from the TCGA data set, no sample belongs to inv(3) subtype with the Papaemmanuil et al. classification.

Furthermore, the Jaccard similarity coefficient was calculated to measure the similarity between each pair of cohorts. It is defined as the size of the intersection divided by the size of the union of the subtype-specific gene sets (Table S3), thus a higher proportion indicates a higher similarity between sample sets. In general, similar proportions are shown for different pairs of cohorts. Subtypes with good survival including CBFB-MYH11, PML-RARA and CEBPA Biallelic generally have higher Jaccard coefficients between the cohorts, ranging from 12% to 20%. The coefficients of poor survival subtypes such as TP53-mutant and MLL are lower. The NPM1 subtype is also distinct from the other subtypes that its Jaccard similarity coefficient between the BeatAML and Clinseq cohorts is the highest (28%), indicating the concordance between the two cohorts. However, the coefficients for the NPM1 group between the TCGA and other cohorts are much lower (8% and 10% for BeatAML and Clinseq, respectively). This suggests a large difference in the NPM1 subtype between the TCGA from the BeatAML and Clinseq cohorts. In addition, to quantify the similarity of gene expression distributions of the identified subtype-specific genes between cohorts, we used the set of top 25 subtype-specific genes to calculate the correlations of median expression for each subtype between BeatAML and other cohorts (Figure S5). The result shows
that the gene expression for the top 25 subtype-specific genes has a strong correlation between BeatAML and other cohorts (Figure S5).

To present an overview of how well the global gene expression data corresponds to the genomic subtypes, we performed the tSNE analysis using: a) all genes and b) the top 25 subtype-specific genes in each subtype of BeatAML cohort. Using top 25 subtype-specific genes, we found a marked increase in the separation between subtypes (Figure 4). Using all genes, only the PML-RARA subtype is well separated (Figure S6). This suggests that most patients had gene expression characteristics consistent with the genomic subtype in the identified subtype-specific genes rather than all genes. We excluded the “other” group from the tSNE analysis to maintain the pure molecular distinct subtypes. The tSNE plot including the “other” group can be found in Figure S7. The tSNE analyses for the top 25 subtype-specific genes identified in the TCGA and Clinseq cohorts are presented in Figure S8A,B, respectively. Similar to results based on the BeatAML cohort, both show a good separation between the subtypes. The color-maps for the top 25 subtype-specific genes identified in individual cohorts are presented in Figures S13–S15. We further use the 729 subtype-specific genes identified in the BeatAML and validated in both the TCGA and Clinseq cohorts from Figure 4 for the tSNE analyses. The results for the BeatAML, Clinseq and TCGA cohorts are presented in Figure S8C–E, respectively. The subtypes are well separated, especially the main subtypes such as NPM1, TP53-mutant, CBFβ-MYH11, PML-RARA, CEBPA<sub>Biallelic</sub> and RUNX1-RUNX1T1.

We further investigated the contribution of the subtype-specific genes to the separation of French-American-British (FAB) subtypes of AML<sup>24</sup> which capture the level of maturation of the cancer cells. This

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**FIGURE 2**  Gene-level expression distribution of PTPRG gene across 11 molecular subtypes in A, BeatAML, B, Clinseq and C, TCGA cohort and protein expression distribution of PTPRG gene across 11 molecular subtypes in D, Clinseq cohort. Gene expression is illustrated in log<sub>2</sub>(TPM + 1) and the protein levels of the gene are logged relative ratios [Color figure can be viewed at wileyonlinelibrary.com]
might indicate whether the subtype-defining mutations introduce a differentiation block and expansion. We selected two largest subgroups NPM1 and TP53-mutant from the BeatAML cohort, collected samples with available FAB information and did the tSNE analysis using the mRNA expression of top 25 subtype-specific genes of the subtypes. For NPM1 subgroup, as shown in Figure S9A, we see a clear separation for FAB subtypes of M1, M2, M4 and M5. However, for TP53-mutant subgroup, some FAB subtypes such as M2 are not well clustered; see Figure S9B. Even though the results are limited due to the few samples with available FAB subtype information, the tSNE analysis suggests that the expression of subtype-specific genes contain information for separation of FAB subtypes.

### 3.3 Subtype-specific protein levels

The same validation process was applied to the proteomics data from the Clinseq cohort. Taking PTPRG gene as an example (Figure 2D), the protein levels of PTPRG gene is highly correlated with gene expressions, as they all have higher values in PML-RARA subtype and lower values in the rest subtypes. Since some genes were not detected in the proteomics data, we included more genes from the subtype-specific gene list and selected the top 25 overlapping genes. So, 41% of these selected genes have high correlation between gene expression and protein levels (with correlation coefficient > 0.5). More intuitively, Figure 3 shows the protein levels of the selected genes for each subtype. Although the validation at protein level is not as clear as at the gene expression level, we still can observe consistent patterns, for example in the NPM1 and PML-RARA subtypes.

### 3.4 Biological pathways

To assess the relations of the identified subtype-specific genes to the biological pathways and processes, we used the Reactome database to analyze the pathway enrichment analysis of the top 25 subtype-
pecific genes in each subtype. The significant pathways (p-value < .05) and statistics for each subtype are listed in Table S4. For example, the most significant pathway (p-value = 0.0019) related to CBFB-MYH11 subtype-specific genes is “RUNX2 regulates osteoblast differentiation”. The RUNX family of transcription factors plays a critical role in hematopoiesis, and the RUNX1 transcription factor is frequently translocated in AML.26 As a RUNX family member, the relation of RUNX2 to CBFB-MYH11 specific genes underlines the fact that RUNX family members can function in complex with CBFB.27 The roles of RUNX2 and CBFB in skeletal development have also been well studied elsewhere.28,29 In addition, the glance of pathway diagram would bring an overview of the connectivity and flow of information in biological systems. Taking the most significant pathway related to CEBPA^{Biallelic} subtype-specific genes (p-value < .001), “Glucuronidation”, for example, we demonstrated the pathway associated with the subtype-specific genes (Figure S10). Four of a total 24 genes of this pathway (16.7%) including UGT2A3, UGT2B10, UGT2B11 and UGT2B28 are represented in 25 top subtype-specific genes of CEBPA^{Biallelic} subtype, indicating the pathway enrichment.

4 | DISCUSSION AND CONCLUSION

We have provided a comprehensive view of the transcriptomic landscape of molecular subtype-specific mRNA expression of AML based on 955 RNA-seq samples from three different cohorts. A sophisticated statistical methodology has been used to identify and validate 729 subtype-specific genes across molecular subtypes. These results suggest that the gene-expression profiles can be used to characterize the molecular subtypes of AML. We have provided a comprehensive data portal which can serve a public resource to be used for hypothesis generation, that is, the discovery of new biomarkers for drug targets.

Rather than discovery of new genomic subtypes, our aim was to provide more characterizations to existing molecular subtypes that have been shown to have strong clinical relevance. Currently in the clinic, the WHO and the ELN classifications have the most impact on the treatment decision, but these are based on biology, morphology and medical history. We had chosen the molecular classification of Papaemmanuil et al. as it provides purely biological distinct subtypes. The value of the molecular classification will increase over time as we get more information on subtype-specific markers and on treatment response. Instead of gene signatures, we provide single gene markers specific to each subtypes. The key advantages are that (i) the single genes may have more biological investigations, and (ii) they are more easily measured in terms of sample requirements, especially if they are validated at single protein level.

The PTPRG gene, a top-ranking gene specific to the PML-RARA subtype in Figure 2, is strongly validated across cohorts and at protein level. This has been reported as a tumor-suppressor gene in not only AML but also other cancers, that is, nasopharyngeal carcinoma.30 So, PML-RARA fusion is a known initiating event for the acute promyelocytic leukemia (APL), and PTPRG mutation was discovered in APL samples.31 Note, PTPRG is a member of the protein tyrosine phosphatase (PTP) family of signaling molecules that are involved in cell cycle, differentiation and oncogenic transformation. It has been identified as a significant gene in childhood acute lymphoblastic leukemia since the phosphatase induces dephosphorylation of ERK which provides a potential therapeutic target for RAS-related leukemias.32 Also, PTPs have been detected in AML previously,33 but not with specificity to any subtypes. The current result suggests a biological role of PTPRG specific to the PML-RARA fusion subtype. Thus, by identifying subtype-specific genes we can discover biomarkers that are specific to each subtype. These genomic biomarkers could have more biological investigations by clinical researchers.

Different molecular subtypes are clearly distinguished from each other in the discovery set (BeatAML) and similar patterns are observed in the validation sets (TCGA and Clinseq), especially for NPM1, CEBPA^{Biallelic}, CBFB-MYH11, PML-RARA and RUNX1-RUNX1T1. This partially agreed with the analysis in,34 where they clustered patients and genes on the basis of similarity of expression distributions by using unsupervised hierarchical clustering analysis and found patients with t(8;21), inv(16) mutations—RUNX1-RUNX1T1 and CBFB-MYH11 fusions—had gene expression patterns specific to t(8;21) and inv(16) subtypes. Among the subtypes, the PML-RARA and NPM1 subtypes have the largest number of subtype-specific genes. Such a large proportion of subtype-specific genes is typically a reflection of a biologically distinct entity. We observe the same phenomenon, for example, if we compare estrogen-receptor (ER)-positive vs ER-negative breast cancers, or lung adenocarcinoma vs squamous-cell carcinoma.

Further explorations could be made based on the identified subtype-specific biomarkers. For example, we have found two subtype-specific genes from CEBPA^{Biallelic} subtype, ZBTB20 and ARHGEF6, in the set of transcription factor (TF) targets of CEBPA.
collected from the Molecular Signatures Database (MSigDB v7.1). The expression distributions (Figures S11 and S12) show the specificity of these genes to CEBPA-Biallelic with a higher expression of the subtype over the rest groups in the BeatAML as well as the validated cohorts. This suggests a role of the CEBPA mutation in the changes of expression of the TF targets.

Our study had some limitations. Because they were based on retrospective samples, the sample collection in the different cohorts was heterogeneous. For example, not all samples across the different cohorts were taken uniformly at the time of diagnosis; also, both bone marrow and peripheral-blood mononuclear cells were in use, identified using Ficoll gradient centrifugation rather than using the CD38+ marker. These effects tend to increase statistical variability, so may reduce the sensitivity in detecting the subtype-specific markers and explain some lack of replicability across the cohorts. However, they do not reduce the specificity of our results.

AUTHOR CONTRIBUTIONS
Trung Nghia Vu and Yudi Pawitan initiated and coordinated the study; Tian Mou performed data analysis; Tian Mou, Yudi Pawitan, Trung Nghia Vu contributed to method development and manuscript writing. Matthias Stah, Mattias Vesterlund, Rozbeh Jafari, Anna Bohlin, Albin Österroos, Loannis Siavelis, Helena Bäckvall, Helena Bäckvall, Tom Erkers, Santeri Kiviluoto, Brinton Seashore-Ludlow, Päivi Östling, Lukas M. Orre, Olli Kallioniemi, Sören Lehmann, Janne Lehtiö also contributed to manuscript writing and the Clinseq cohort including the acquisition and processing of patient samples, and the collection of clinical data. Wenjiang Deng and Trung Nghia Vu contributed to the pre-processing of RNA sequencing data. All authors read and approved the final manuscript.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT
The results of subtype-specific genes discovery in this study can be found at: https://nghiavtr.shinyapps.io/AMLSubtypeSpecificDiscovery/.

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**SUPPORTING INFORMATION**

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

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