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BRIEF REPORT



CD56, HLA-DR, and CD45 recognize a subtype of childhood AML harboring CBFA2T3-GLIS2 fusion transcript

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Abstract

The presence of CBFA2T3-GLIS2 fusion gene has been identified in childhood Acute Myeloid Leukemia (AML). In view of the genomic studies indicating a distinct gene expression profile, we evaluated the role of immunophenotyping in characterizing a rare subtype of AML-CBFA2T3-GLIS2 rearranged. Immunophenotypic data were obtained by studying a cohort of 20 pediatric CBFA2T3-GLIS2-AML and 77 AML patients not carrying the fusion transcript. Enrolled cases were included in the Associazione Italiana di Ematologia Oncologia Pediatrica (AIEOP) AML trials and immunophenotypes were compared using different statistical approaches. By multiple computational procedures, we identified two main core antigens responsible for the identification of the CBFA2T3-GLIS2-AML. CD56 showed the highest performance in single marker evaluation (AUC = 0.89) and granted the most accurate prediction when used in combination with HLA-DR (AUC = 0.97) displaying a 93% sensitivity and 99% specificity. We also observed a weak-to-negative CD45 expression, being exceptional in AML. We here provide evidence that the combination of HLA-DR negativity and intense bright CD56 expression detects a rare and aggressive pediatric AML genetic lesion improving the diagnosis performance.

KEYWORDS

acute myeloid leukemia, CBFA2T3-GLIS2, computational analysis, immunophenotyping

Giuseppe Basso and Barbara Buldini contributed equally to this work as last co-authors. [†]deceased.

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1 | INTRODUCTION

A recurrent cryptic inversion of chromosome 16 [inv (16) (p13.3q24.3)] fuses a portion of CBFA2T3 gene in frame with a region of GLIS2 gene encoding for CBFA2T3-GLIS2 fusion transcript [1,2]. This chimeric oncogene accounts for 18% of pediatric non-Down syndrome acute megakaryoblastic leukemia (non-DS-AMKL) patients and 8% of pediatric cytogenetically normal AML (CN-AML) associated with different French-American-British (FAB) subtypes [1-3]. This genetic lesion confers a particularly poor prognosis, with event-free survival (EFS) ranging between 8% and 33% in different studies [1-5]. The gene expression profile of CBFA2T3-GLIS2 AML has been associated with up-regulation of both Hedgehog (HH) and bone morphogenic protein (BMP) signaling [2,6,7]. In view of the genomic studies indicating a distinct gene expression profile compared with other genetic subtypes of childhood AML [2], we evaluated the immunophenotypic profile of pediatric AML CBFA2T3-GLIS2 rearranged.

TABLE 1Clinical characteristics fornon-DS and non-APL AML patients(CBFA2T3-GLIS2 positive and negative)enrolled in the study

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2 | MATERIALS AND METHODS

From April 2001 to March 2017, we identified 25 CBFA2T3-GLIS2positive cases. Five samples were excluded from this analysis due to missing immunophenotypic data. The immunophenotype of 20 CBFA2T3-GLIS2-positive cases was compared with a cohort of 77 CBFA2T3-GLIS2-negative AML patients, including also two cases of Down syndrome myeloid leukemia (ML-DS) and nine cases of acute promyelocytic leukemia (APL), collected in the same time interval. All patients were enrolled in the Associazione Italiana di Ematologia Oncologia Pediatrica (AIEOP) AML trials (AIEOP AML 2002, AIEOP ICC APL Study 01). The biological material used in this study was centralized at the Pediatric Hematology Oncology reference AIEOP Laboratory of Padua. Clinical features are detailed in Table 1, data on molecular genetics in Table S1. Written informed consent to the use of the patient's data and biological material was obtained from parents/legal guardians in accordance with the Declaration of Helsinki. Local institutional ethical committees approved the study, including

	CBFA2T3-GLIS2 positive		CBFA2T3-GLIS2 negative		n
	N	%	N	%	٢
Total cohort	18	100	66	100	
Age (years)					
<10	18	100	36	54.5	<0.01
>10	0	0	30	45.5	<0.01
Gender					
Female	10	55.6	37	56.1	0.11
Male	8	44.4	29	43.9	0.11
WBC					
≤10,000	3	16.7	23	34.8	0.16
10,000 ≤ 100,000	15	83.3	37	56.1	0.05
>100,000	0	0	6	9.1	0.33
FAB					
M0	0	0	2	3	1
M1	1	5.5	6	9.1	1
M2	1	5.5	20	27.4	0.06
M2Eo	0	0	2	3	1
M4	0	0	17	25.8	0.02
M4Eo	0	0	1	1.5	1
M5	1	5.5	11	16.7	0.44
M6	0	0	1	1.5	1
M7	14	78.0	3	4.5	<0.01
Other	1	5.5	5	7.5	1
BM blasts (median %)		50		64	0.37
Extramedullary involvement ^a	4	22.2	6	9.1	0.21

Note: p values have been calculated using Fisher's statistics. The bold values denote statistical significance at the p < 0.05 level.

^aOne patient with additional CNS involvement, three patients without CNS involvement.

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the collection of bone marrow samples (BM) for diagnostic work-up, along with the national protocols. Diagnosis of *de novo* AML was performed according to WHO classification criteria [8,9]. Samples were screened for the presence of t(8;21), inv(16), t(16;16), t(15;17) with associated molecular transcripts, namely core-binding factor (CBF)- β abnormalities and PML-RAR α ; moreover, KMT2A rearrangements [9,10], Internal Tandem Duplication of FLT3 mutation (FLT3-ITD) [11]

and other rare mutations were screened (Table S1).

2.1 | Flow cytometry

Initial evaluation of qualitative results from the whole immunophenotypic characterization was performed at diagnosis according to WHO criteria and the latest consensus guidelines for flow-cytometry immunophenotyping published by the AIEOP-BFM working group [12]. Quantitative values of mean fluorescence intensity (MFI) was retrieved for 14 antigens: CD45 ECD, HLA-DR PE-Cv7. CD56 PE, CD11a PE, CD11b PE, CD13 FITC, CD15 FITC, CD41 FITC, CD61 PE-Cy7, CD38 PE-Cy5, Myeloperoxidase (MPO) FITC, CD33 PE-Cy5, CD34 PE-Cy5, and CD117 PE. Samples were processed and analyzed in the Hemato Oncology Laboratory of Padua, according to standardized operating procedures (SOP) previously described [8,12]. Flow cytometry methods have been detailed in a Supplementary document according to MIFlowCvt standards.

2.2 | Statistical analyses

Statistical procedures were performed to identify putative antigens correlated with the presence of CBFA2T3-GLIS2 mutation and estimate their accuracy in classifying patients as "GLIS" (with mutation) or "nGLIS" (without mutation) samples. All statistical analyses were performed using R software [13]: Random Forest analyses including Mean Decrease in Accuracy (MDA) and Mean Decrease in Gini coefficient (MDG) procedures were performed using "randomForest" package [14]; multiple testing corrections were evaluated with "multtest" package [15]; permutation tests were retrieved using k-Nearest Neighbor classifier and "caret" package; classification model's performance was evaluated using Area Under Receiver Operating Characteristics (AUROC or AUC) calculated with "ROCR" package; principal component analysis (PCA) was obtained using "factoextra" and "factoMiner" packages. Figure 1A summarizes the analysis steps performed in the whole study starting from cohort description through feature selection, classification, and performance, ending to spatial visualization of the results.

3 | RESULTS AND DISCUSSION

Ninety-seven patients were investigated for antigen (N = 14) associations using different computational analyses (Figure 1A). First, supervised random forests analysis was performed to measure the

importance of all variables and their ability in classifying the data appropriately [14]. MDA reflects the impact of each antigen on the accuracy of the prediction, model-antigens with a large mean decrease in accuracy, being more important for classification. As shown in Figure 1B, CD56 and HLA-DR antigens were the most accurate markers for the identification of "GLIS" AML cases in MDA evaluation; MDG (Figure 1C) confirmed MDA results, as detailed in Table S2. The same markers were identified by antigen selection



FIGURE 1 Variable selection procedures and data presentation. (A) Analysis steps performed from cohort description to spatial visualization of the results. (B) Mean decrease in accuracy (MDA) results using "randomForest" package in R software. CD56 and HLA-DR have the largest mean decrease scores and are considered the most accurate antigens for classification of the data. (C) Mean decrease in Gini (MDG) coefficient confirms CD56 and HLA-DR antigens as the most accurate markers in separating "GLIS" and "nGLIS" patients. (D) Principal component analysis (PCA) results using "factoextra" and "factoMiner" packages. "GLIS" group (in blue) with HLA-DR [Color figure can be viewed at wileyonlinelibrary.com]

based on Wilcoxon's test and Bonferroni procedure for multiple testing corrections [15]. As shown in Table S3, CD56 (p = 0.00014) and HLA-DR (p = 0.00014) were the most discriminating markers followed by CD45 (p = 0.00686), MPO (p = 0.00882), CD38 (p = 0.01932), and CD11a (p = 0.02282). To test the classification strength of identified markers, for each of the 10,000 permutations we randomly assigned patients either to training (total N = 64: N = 13 "GLIS", N = 51"nGLIS") or validation (total N = 33: N = 7 "GLIS," N = 26 "nGLIS") sets. We further calculated the performance of the two markers and their combination in classifying "GLIS" and "nGLIS" patients using ROC curves. Weighted mean (WM) was calculated for accuracy measures including sensitivity, specificity, PPV (positive predictive value), and NPV (negative predictive value) as well as for AUC permutations. CD56 held the highest score in single-marker model evaluation (AUC = 0.89) followed by HLA-DR (AUC = 0.86): remarkably, their combination granted the most accurate model prediction with AUC = 0.97, as well as the highest weighted mean sensitivity (0.93), specificity (0.99), PPV (0.96), and NPV (0.98), Biplot (PCA) was performed on all patients using only CD56 and HLA-DR to clearly show the correlation between groups and antigens supporting the strength of the selected signature (Figure 1D): "GLIS" group was strongly associated with expression of CD56, while "nGLIS" correlated mainly with HLA-DR antigen expression (details in Table S4).

The aim of this study was to evaluate the role of multiparametric flow cytometry analysis in the identification of CBFA2T3-GLIS2 positive AML as previously reported for other genetic subtypes. Our results show that the combination of low HLA-DR together with CD56 as intensive bright provides the highest accuracy in classifying an AML case as "GLIS." The lack of HLA-DR is one of the most sensitive feature typical of APL with respect to other AML subtypes [16] and is described as negative also in some NPM1-mutated AMLs [17-19]. Antibodies against CD56 are primarily used to detect natural killer (NK) cells but are also useful in the evaluation of pediatric solid tumors, such as neuroendocrine and neuroectodermal malignancies (e.g., neuroblastoma, primitive neuroectodermal tumor) [20, 21]. We attempted a CD56 intensity comparison in GLIS-AML and a solid tumor finding a comparable CD56 bright expression (Figures 2 and S1A,B). In AML, CD56 expression has been frequently associated with t(8;21) RUNX1-RUNX1T1 [22] but with a lower intensity than compared with "GLIS" cases. Thiollier et al. previously showed that CBFA2T3-GLIS2+ AMKL exhibits a distinct gene-expression signature: surface marker NCAM1 (CD56) presented a mean differential expression of 35-fold by microarray and of more than 200-fold by RNA-seq with respect to other non-DS AMKL. Using flow-cytometry, they also observed that AMKL blasts carrying CBFA2T3-GLIS2 fusion were CD41+/CD56 + [6]. In our cohort, we observed by flow cytometry a CD56 median differential expression of 29-fold with respect to AML nGLIS (Figure S1B). Moreover, in diagnostic immunophenotyping either lack of or low expression of CD45 is uncommon in AML (Figure S1A); negativity for CD45 associated with positivity for CD56 could be suggestive of non-hematological malignancies [20,21]. In the group of "GLIS" patients here described, CD45 was under-expressed with respect to other AML subtypes (not statistically relevant).



FIGURE 2 Flow cytometry analyses of BM samples. CBFA2T3-GLIS2 positive AML immunophenotype (Case 1) may be misinterpreted with non-hematopoietic tumors BM involvement (Case 2) due to the comparable high expression of CD56 and downregulation of CD45 [Color figure can be viewed at wileyonlinelibrary.com]

The peculiar immunophenotype we discuss also matches the RAM phenotype described by Eidenschink Brodersen et al. [23] The RAM phenotype was characterized by bright CD56, dim-to-negative CD45 and CD38, lack of HLA-DR; it was primarily restricted to very young patients and associated with an extremely poor outcome [23,24]. Pardo et al., investigated the relationship between CD56 expression and prognosis in AML patients enrolled in the Children's Oncology Group Trial AAML0531 [25]. In particular, their Cohort 3 was characterized by a unique immunophenotype (RAM phenotype) and a high prevalence of patients with CBFA2T3-GLIS2 fusion transcript. In our study, we corroborate a highly specific immunophenotype for this fusion transcript in an independent cohort of patients enrolled in the AIEOP study. Analysis of our patients' characteristics confirmed that the CBFA2T3-GLIS2 rearrangement is typical of very young children [26] and is not restricted to M7 subtype (Table 1) [1]. Although, despite this characteristic immunophenotype, no morphological aspects were 848

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suggestive for "GLIS": BM smears could also mimic BM invasion by non-hematopoietic tumors (Figure 3A–D). Clinical features at leukemia presentation could include extramedullary involvement: frequency of extramedullary leukemia in our CBFA2T3-GLIS2 patients was higher (22.2%, Table 1) than that described in pediatric AML [27]. Notably, in two of four cases, extramedullary involvement (including massive cranial bone, ribs, and lumbosacral column involvement, respectively) primarily raised the suspicion of non-hematopoietic tumor instead of acute leukemia.

Accurate diagnosis of childhood hematological malignancies is often challenging and based on the combination of morphology, flow cytometry, cytogenetics, and molecular biology data. Here, we describe an immunophenotypic profile specific for a rare cytogenetic lesion (*CBFA2T3-GLIS2*), clearly discriminated from other AML subtypes and characterized by weak-to-negative CD45 expression unusual in AML, HLA-DR negativity (a finding uncommon in AML apart from APL and NPM1-mutated cases), and intense bright CD56 expression (more suggestive of non-hematological malignancies). This immunophenotypic profile not suggestive of AML could be associated with clinical presentation and BM morphologic features that overlap non-hematopoietic tumors as shown in some of our patients.

In the presence of symptoms related to organ failure or extramedullary involvement, in young children with no clear diagnosis

of solid tumors, we propose to add a BM aspirate to the diagnostic workup, even if peripheral blood count does not portend leukemia diagnosis. Whenever suspicious tumor cells are detected, further comprehensive phenotypic characterization must be performed. Immunophenotyping by flow cytometry is mandatory for a correct AML diagnosis and can provide the indication of molecular biology before other invasive diagnostic procedures are planned.

Overall, our study supports a specific immunophenotype highly suggestive of the AML subtype carrying *CBFA2T3-GLIS2* rearrangement. Our results indicate that an accurate flow cytometry evaluation of leukemia cells at diagnosis allows a rapid identification of this AML subtype even when AML morphological aspect and clinical presentation may not be suggestive of this rare and aggressive pediatric AML entity.

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FIGURE 3 Examples of morphological presentation of bone marrow aspirates in CBFA2T3-GLIS2 positive patients. (A) Bone marrow aspirate showing AML infiltration overlapping BM involvement by non-hematopoietic tumors; (B) High-power magnification of leukemia cells clump; (C) infant with clinical presentation suggestive of non-hematopoietic tumor: in normal regenerating BM aspirates, isolated cells of medium size with large nuclei and prominent nucleoli, basophilic cytoplasm with blebs, no granules or Auer rods may be observed; (D) Leukemia cells suggestive of FAB-M7 morphology [Color figure can be viewed at wileyonlinelibrary.com]

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AUTHOR CONTRIBUTIONS

Andrea Zangrando: Conceptualization; data curation; formal analysis; investigation; methodology; software; validation; visualization; writing-original draft; writing-review & editing. Francesca Cavagnero: Formal analysis; investigation; methodology. Pamela Scarparo: Data curation; formal analysis; investigation; visualization. Elena Varotto: Data curation; formal analysis; investigation; visualization; writingreview & editing. Samuela Francescato: Investigation; visualization. Claudia Tregnago: Data curation. Rosanna Cuccurullo: Resources. Franca Fagioli: Resources. Luca Lo Nigro: Resources. Riccardo Masetti: Resources. Maria Caterina Putti: Resources. Carmelo Rizzari: Resources. Nicola Santoro: Resources. Andrea Pession: Data curation; resources. Martina Pigazzi: Conceptualization; data curation; funding acquisition; investigation; methodology; writing-original draft; writing-review & editing. Franco Locatelli: Conceptualization; data curation; methodology; resources; supervision; writing-original draft. Giuseppo Basso: Conceptualization; funding acquisition; methodology; project administration; resources; supervision; writing-original draft; writing-review & editing. Barbara Buldini: Conceptualization; data curation; investigation; methodology; project administration; resources; supervision; validation; writing-original draft; writingreview & editing.

CONFLICT OF INTEREST

The authors declare no competing financial interests.

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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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