

Letter

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Impaired Overall Survival in Young Patients With Acute Myeloid Leukemia and Variants in Genes Predisposing for Myeloid Malignancies

Martin Kirschner^{1,2}, Benjamin Rolles^{1,2}, Martina Crysandt^{1,2}, Christoph Röllig³, Friedrich Stölzel³, Michael Kramer³, Martin Bornhäuser³, Hubert Serve⁴, Uwe Platzbecker⁵, Carsten Müller-Tidow⁶, Kim Kricheldorf^{1,2}, Margherita Vieri^{1,2}, Matthias Begemann⁷, Angela Maurer^{1,2}, Marcin W. Wlodarski^{8,9}, Sushree S. Sahoo⁹, Tim H. Brummendorf^{1,2}, Edgar Jost^{1,2}, Fabian Beier^{1,2}

Correspondence: Martin Kirschner (mkirschner@ukaachen.de).

Acute myeloid leukemia (AML) is an aggressive myeloid neoplasm with a relevant subgroup evolving from inherited disorders. According to the WHO 2016 classification, the latter group is categorized as “Myeloid neoplasms with germ line predisposition (MNGLP),”¹ comprising various syndromes based on germline mutations in genes such as *CEBPA*, *GATA2*, *RUNX1*, or *SAMD9*² as well as bone marrow failure syndromes such as telomere biology disorders (TBD).¹ The increased risk of AML development at younger age is one common criteria of this category.²

Our study focused on the incidence of TBD as a subcohort of MNGLP in younger AML patients with aberrant karyotype. TBD patients are at particularly high risk of malignant transformation both toward solid tumors and hematologic neoplasms, with the risk of MDS and AML development increased up to 2700- and 200-fold, respectively.³ The identification of classical

TBD such as dyskeratosis congenita (DKC) is based on family history and the typical clinical triad (leukoplakia, nail dystrophy, abnormal skin coloring) mostly predominant in younger patients. Due to a less specific and more heterogeneous spectrum of phenotypes in adult-onset TBD, classical DKC signs are often missing and consequently, accurate diagnosis can be challenging. This together with an overall limited awareness of late-onset genetic disorders with first manifestation in adult age results in significant underdiagnosis.⁴ As a result, adult AML may often be the first manifestation of TBD in selected cases.^{2,5}

TBDs are characterized by impaired telomere maintenance eventually leading to accelerated and functionally critical telomere shortening which in return is associated with chromosomal instability.³ Therefore, AML development in TBD is supposed to be driven mostly by chromosomal aberrations resulting from telomere-mediated chromosomal fusion events or aneuploidy.⁶ In line with this model, AML arising from TBD is supposed to go along with an increased frequency of aberrant karyotypes probably predominantly involving chromosome arms with short telomeres.⁴ While the risk of AML development in TBD is known, the reverse incidence of an underlying TBD in adult patients with AML is unclear to date.³

Based on these considerations, the present study aimed to determine the incidence of underlying TBD cases in young newly diagnosed AML patients with aberrant karyotype. Telomere length (TL) screening via PCR in nonclonal cells was performed in remission samples following induction therapy to investigate the relationship between the degree of preexistent telomere shortening and onset of AML. In order not to miss other additional MNGLP, we performed a comprehensive genetic screening for non-TBD-MNGLP in this preselected cohort.

The database of the German Study Alliance Leukemia (SAL) registry including 5207 patients with AML was screened for patients below 35 years ($n = 577$) fulfilling the following criteria: (1) blast-free state/remission after chemotherapy, (2) aberrant karyotype (≥ 3 aberrations) detected in diagnostic karyotype or FISH analysis, and (3) available samples of peripheral blood or bone marrow (Figure 1A). Detailed methods are described in the Suppl. section.

Using the screening approach mentioned above, we were able to identify 29 patients with DNA for next generation sequencing (NGS) analysis and available data for overall survival (OS) in 25 patients and. All patients have been classified as de novo AML by the treating physicians (for detailed characteristics, see Table 1).

¹Department of Hematology, Oncology, Hemostaseology and Stem Cell Transplantation, Medical Faculty, RWTH Aachen University, Germany

²Center for Integrated Oncology Aachen Bonn Cologne Düsseldorf (CIO ABCD), Germany

³Department of Medicine I, University Hospital Carl Gustav Carus Dresden, Technical University Dresden, Germany

⁴Department of Medicine II, Hematology/Oncology, Goethe University, Frankfurt/Main, Germany

⁵Department of Medicine I, Hematology, Cellular Therapy, Hemostaseology, University of Leipzig, Germany

⁶Department of Medicine V, Hematology, Oncology and Rheumatology, University Hospital Heidelberg, Germany

⁷Institute of Human Genetics, Medical Faculty, RWTH Aachen University, Germany

⁸Department of Pediatrics and Adolescent Medicine, Division of Pediatric Hematology and Oncology, Faculty of Medicine, Medical Center, University of Freiburg, Germany.

⁹Department of Hematology, St. Jude Children's Research Hospital, Memphis, TN, USA

THB, EJ, and FB have contributed equally to this work.

Supplemental digital content is available for this article.

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HemaSphere (2022) 6:11(e787).

<http://dx.doi.org/10.1097/HS9.0000000000000787>.

Received: May 29, 2022 / Accepted: September 4, 2022

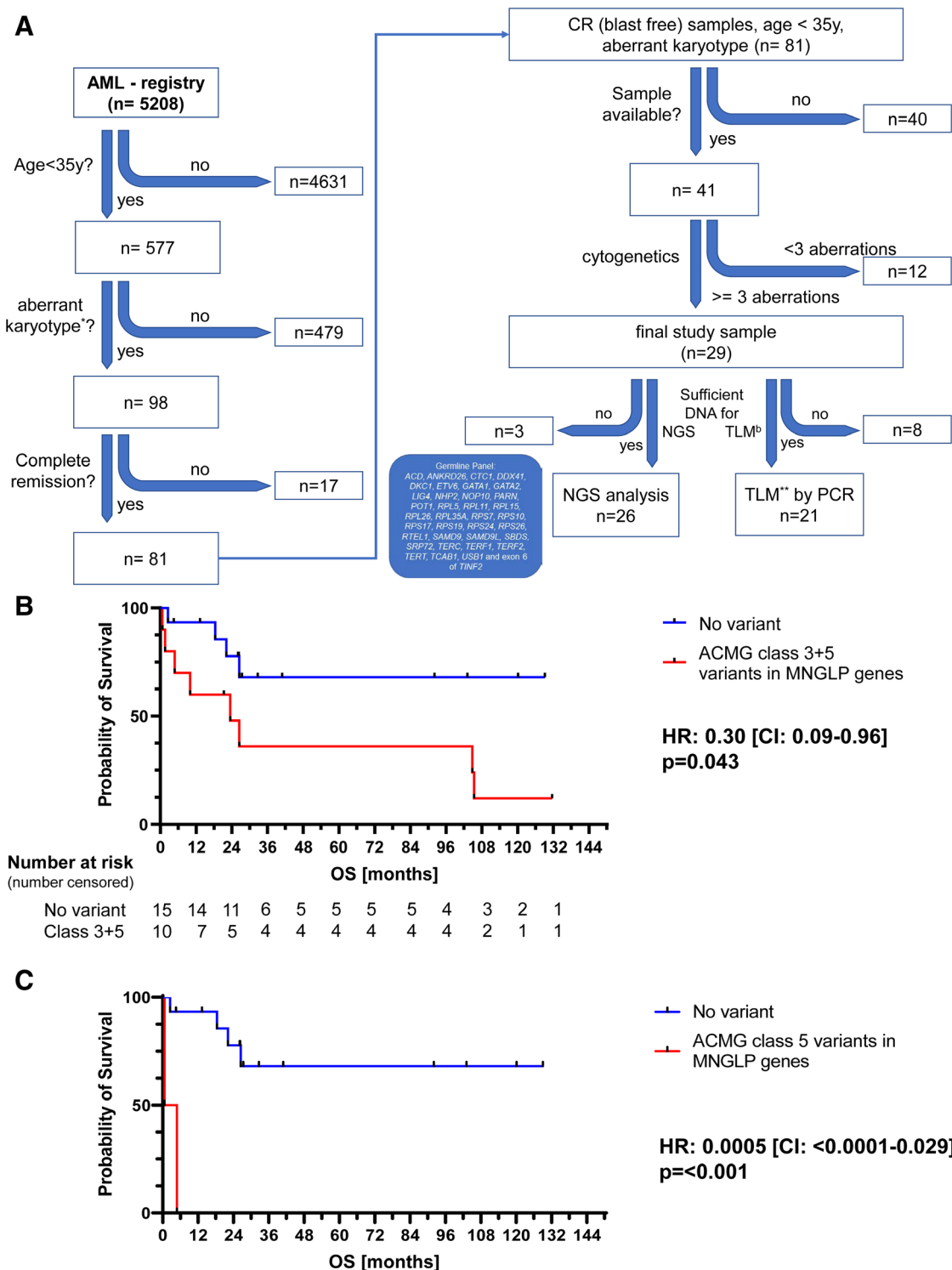


Figure 1. Flow chart of sample availability (A) and overall survival of patients with non-MNGLP (n = 15) and with variants for MNGLP (n = 10) (B) or pathogenic mutations for MNGLP (n = 2) (C) is shown. (A) * = complex aberrant or trisomy 8 according to register log; ** = telomere length measurement (TLM). (B+C) Patients are not censored for allogeneic transplantation due to the small number of patients. For three patients no molecular analysis data were available and for 1 patient no overall survival was documented (cases #22, #25, #26, and #29, excluded in this analysis). Median follow-up was 26.3 months [Q1–Q3: 13.3–92.04]. Log Rank Test was used for statistical analysis. MNGLP = myeloid neoplasms with germ line predisposition.

First, we analyzed whether TL prescreening in remission samples can be used to “enrich” for patients with underlying TBD (n = 21, see Supplement for details). Mean TL of all patients was not significantly shortened compared to age-adjusted controls in line with previous data.⁸ However, using the 1% percentile as

cutoff, we found 6 patients with significantly shortened TL. NGS analysis of this subcohort revealed two variants of unknown significance (VUS) in the TBD-associated genes *TINF2* and *RPA1* according to the American College of medical genetics (Table 1). In order not to miss a TBD or other MNGLP case, we expanded

Table 1.
Summary of Molecular and Clinical Data of All 29 Patients From the SAL-AML Registry

Patient No, n = 29	Sex	Age	Cytogenetics (≥3 Aberrations)	Allogeneic SCT	CR After Induction	AML-Relapse	Death	Affected Genes Hereditary (ACMG class)	ACMG Criteria (Detail)	Affected Genes Somatic; Pathogenic Variants	TL Below 1st Percentile	OS (mo)
1	M	33	46,XY [7]; 46,XY(9)(p11;p24), t(1;12)(p34;p13), del(12)(p11-12) [23]	Yes	Yes	No	No	/	/	/		92.04
2	M	24	46,XY [3]; 46,XY,del(7)(q22), inc [1]; 46,XY,-6, del(7)(q22), inv(16)(p13q22), +mar, inc [1]	No	Yes	Yes	Yes	TINF2 c.62A>G; p.(Gln21Arg) (class 3)	PM1, PP4			104.73
3	F	22	50,XX,add(2)(p7), del(3)(p14), der(3)(p711), +,-6mar, inc [cp10]; 46 chromosomes [5]	Yes	Yes	No	Yes	SBD5 c.258 + 27>C/ c.183_184delinsCT; p.(Lys62Ter) (class 5)	PVS1, PM2, PP3, PP5	TP 53, two variants (NAF 6% and 78%)		4.80
4	F	26	45,XX,t(9;15)(q17;q1?), der(7)(t10)(13qter-->13q?::10p11-10q22::13q-->13a), t(11;17)(q23;q21), der(11)(t10;11)(q22;p15), del(12)(q22;p15), del(12)	Yes	Yes	Yes	Yes	ANKRD26 c.4924A>G, p.(Asn1642Asp) (class 3)	PM1, PM2, BP4			9.96
5	F	30	46,XX [6]; 47,XX, +22, inv(16)(p13q22), ?del(16)(q22q24) [cp20]	No	Yes	No	No	/	/			103.12
6	F	19	49,XX,+4,t(6;9)(p23;q34), +13,+22[17] FISH: 40% monosomy	Yes	Yes	Yes	Yes	RPA1 c.1397C>G; p.(Ala466Gly) (class 3)	PM1		Yes	105.33
7	F	24	46,XX,t(11;17)(q23;q25)[7]; 48, idem, +i(8)(q10)x2[9]; 47, idem, +8[4]	Yes	Yes	No	No	/	/			129.06
8	M	33	46,XY,add(4)(p16), add(6)(p21), del(12)(q13)[8]; 46,XY, idem, del(7)(p13)[18]; 46,XY[3]; 46,XY,t(4;6)(p16;p11), t(12;13)(p11;q1)	Yes	Yes	No	No	RPA1 c.460G>A; p.(Thr154Ala) (class 3)	PM1, PM2			131.56
9	M	33	48,XY, +8, t(1;19)(q23;p13), +21[2]; 49,XY, +8, t(1;19)(q23;p13), +der(19)t(1;19)(q23;p13), +21[7]; 46,XY[1]	No	No	No	Yes	/	/			2.50
10	F	22	46,XX,der(4)(del(4)(q11)(4;?1)(q11;q23q23), der(7)(del(7)(q11)t(7;?10)(q11;p12), +8, der(10;11)(10qter-->10p12::11q23-->11q23:48,XX,+X,+4,der(5)t(5;6)(q37;?), t(10;11)(p15;q13)[4]	Yes	Yes	Yes	Yes	/	/	ASXL1 NAF 17%; FLT3 NAF 12%	Yes	26.46
11	F	27	63~91<4n>, XXXX,+X,+4, +4, der(5)t(5;6)(q37;?)x2,-5,-6, t(10;11)(p15;q13)	Yes	Yes	No	No	/	/			120.12
12	M	20	46,XY,der(5)(del(5)(q31)(4;5)(?;3), ish: der(5)(EGR1-), ish t(11;?) (5'MLL); 46,XY, der(5)(del(5)(q31)(4;5)(?;q31), ish der(5)(EGR1-)	No	No	No	Yes	TINF2 c.734C>A; p.(Ser245Tyr) (class 3)	PS5, PM1, BS2, BP4, BP6		Yes	1.61
13	M	26	47,XY,t(7)+mar[7]; 48,XY,t(7)+mar,+16[9]; 46,XY[1]	Yes	Yes	Yes	Yes	/	/	PTPN11 (NAF 39%)		22.12
14	F	27	47,XX,t(7;11)(p15;p15), +8,+19[25]	Yes	Yes	No	No	/	/			27.37
15	M	20	43-47,XY,-2,der(4)-5, der(5)-6,-7, der(10), der(12), -13,der(14),add(14)(p), t(15;18)(q10;q10), der(17), -18,+22,+2-5mar	Yes	Yes	Yes	Yes	CTC1 c.2506T>C, p.(Ser836Pro) (class 3)	PM2			26.49
16	M	24	46,XY/ 46,XY,del(8),der(13)(t13;15), der(20)(t8;20), der(21)(t15;21)/ 92,XXX, del(8)x2, der(13)(t13;15)x2, der(20)(t8;20)x2, der(21)(t15;21)x2	Yes	Yes	Yes	Yes	/	/			18.44
17	M	32	46,XY,t(8;21)(q22;q22), t(19;22)(q13;q11.2), ish t(19;22)(wcp22+;wcp22+BCR+)[22]/ 45,X,-Y,t(8;21)(q22;q22), t(19;22)(q13;q11.2)[3]	No	Yes	No	No	/	/			13.31
18*	M	22	48,XX,+8, inv(16)(p13q22), +21	No	Yes	No	No	/	/	KRAS (NAF 63%)		40.88
19	M	32	45,XY,der(6)t(6;8)(q27;p22), der(8)t(8;11)(p22;p12), del(11)(p12p15), dlc(12;16)(p11;q24) [2]/ 46,XY[20]	Yes	Yes	No	No	/	/			26.13
20	M	32	45,X,-Y,del(8)(q13), der(9) (9p13->9q21::8q13->8q22::21q22>21qter), der(14) ins(14;9) (q24::q22;q34), der(21) t(8;21)(q22;q22) t(8;9)(qq24;p13)[19]	No	Yes	No	No	/	/			4.53

(Continued)

Table 1. (Continued)

Patient No, n = 29	Sex	Age	Cytogenetics (≥3 Aberrations)		Allogeneic SCT	CR After Induction	AML-Relapse	Death	Affected Genes Hereditary (ACMG class)	ACMG Criteria (Detail)	Affected Genes Somatic; Pathogenic Variants	TL Below 1st Percentile	OS (mo)
21	F	29	45,XX,add(8)(p11.2),-1,-1,-14,-16,+21,+mar[6]/46,sl,+add(8)(p11.1)[18] 46,XX[1]	Yes	Yes	Yes	No	No	/	/	/	Yes	26.29
22	F	18	41~42,XX,-4,-5,-6,-7,-13,add(17)(p11.1),-18,+2mar[cp3],nuc ish	No	No	No	No	No	/	/	TP53 (VAF 33%)	Yes	Unknown
23	F	31	45,X,-X,t(8;21)(q22;q22),del(9)(q13),del(13)(q12q14)[16]/46,XX[1]	No	Yes	Yes	No	No	SAMD9 c.2642A>G; p.(Asp881Gly) (class 3)	PM1, BS1	DNMT3A (VAF 14%)	Yes	21.33
24	F	30	47,XX,+8,add(13)(q7),add(20)(q7)[15] 46,XX[7]	Yes	Yes	Yes	Yes	No	/	/	/	Yes	32.67
25	F	19	47,XX,t(1;3)(p36;q26),t(8;18)(q24;q11.2),+mar[10]/47,XX[10]	Yes	Yes	Yes	No	No	NA	NA	/	Yes	22.54
26*	F	32	46,XX,t(8;21)(q22;q22),del(9)(q22) 45,X-X,add(2)(q7),t(8;21),del(9)(q7)	Yes	Yes	Yes	No	No	NA	NA	/	Yes	148.78
27	F	28	47,XX,+del(1)(p13),t(9;10;11)(11qter->11q23;9p21->9qter; 9pter->9p21::11q23->11q13::10p13->10qter; 11pter->11q13::10p13->10qter	Yes	Yes	Yes	Yes	Yes	SAMD9 c.2642A>G; p.(Asp881Gly) (class 3)	PM1, BS1	ASXL1 (VAF 42%); NPAS (VAF 44%)	Yes	23.47
28	M	25	46,XY,+1,der(1;7)(q10;q10),inv(9)(p12q13)[25]	No	No	No	No	Yes	GATA2 c.1009C>T; p.(Arg337Ter) (class 5)	PVS1, PM2, PP3, PP5	/	0.62	5.98
29*	M	27	46,XY,der(2)t(2;8)(q35;q22)x2,t(8;15;21)(q22;q21;q22)	No	Yes	Yes	No	Yes	NA	NA	/	Yes	5.98

Bold text highlights mutations in the 2 patients with class 5 variants (confirmed pathogenic).

All patients fulfilled the inclusion criteria: age below 35, confirmed AML as the main diagnosis, aberrant karyotype (≥3 aberrations), Cases 18, 26, and 29 (*) showed a favorable cytogenetic risk, whereas all other samples are classified as adverse risk. Variants were classified according to the ACMG classification and for used abbreviations refer to Richards et al (2015).⁷

ACMG = American College of medical genetics; NA = not available; OS = overall survival; TL = telomere length.

the NGS screening to the remaining 23 patients. We detected 6 additional patients with VUS and 2 with pathogenic mutations in TBD- or other MNGLP-related genes (Table 1) increasing the total number of identified class 3 and 5 variants in our cohort to 34.5% (n = 10/29). In addition to the 2 cases mentioned above, we found 3 additional variants in genes associated with TBD (*CTC1*, *RPA1*, and *TINF2*). For non-TBD-related MNGLP, we found 3 cases with VUS in *SAMD9* (2×) and *ANKRD26* and 2 variants in *GATA2* and *SBD5* previously described as pathogenic (class 5, see Table 1).^{9,10}

Finally, we compared the clinical outcome in patients with variants in MNGLP-associated genes (n = 10) with patients harboring no variants ("non-MNGLP" patients, n = 15). Patients with variants in MNGLP-associated genes showed a significantly reduced OS time compared to non-MNGLP patients (HR: 0.30; 95% CI 0.09-0.96, *P* = 0.043; Figure 1B). Patients with pathogenic variants only (n = 2) showed an even more pronounced difference in OS compared to non-MNGLP patients (HR: 0.0005; 95% CI: <0.0001 to 0.029, *P* ≤ 0.001; Figure 1C). Relapse as a possible cause for impaired survival was reported in 50% (5/10) of our patients with detected variants. Due to the retrospective nature of the cohort, data on the causes of death were unfortunately not available.

In summary, we analyzed the proportion of patients with underlying MNGLP with a particular focus on TBD in young AML patients with aberrant karyotype. Indeed, TL prescreening was able to narrow down the cohort to 6 patients with shortened TL of whom 2 patients were eventually found to have variants in TBD-associated genes. However, it was not able to identify the additional 3 cases identified with variants in TBD-associated genes. One reason for this limitation possibly resides in the impaired diagnostic value of telomere PCR compared to the gold standard diagnostic method flow-FISH (requiring living cells).⁷ Obviously, due to the relatively small sample size of this clinically preselected cohort, larger studies including functional testing are needed to substantiate the false positive/negative test ratio and the final diagnostic value of TL prescreening in this setting.

Few data are available about the incidence of germline predisposition in AML in general. Two recent studies in predominantly older AML patients showed a different distribution pattern of the detected variants^{11,12} with *CHEK2* and *DDX41* as the most frequent variants. In contrast, we predominantly found variants in MNGLP genes associated with bone marrow failure syndromes. This difference can be explained by the different approach in patient selection with the focus on younger patients with chromosomal aberrations and the methods used for variant detection using whole-exome sequencing versus panel based NGS.

Interestingly, we observed a significantly impaired OS in the patients with variants in MNGLP genes compared to the rest of the cohort. This is somewhat in contrast to previous data showing a more favorable outcome at least for patients with *DDX41* mutations.¹³

We were not able to provide detailed analysis for the causes of death in our MNGLP cohort. Patients with bone marrow failure associated MNGLP have in general an increased risk for treatment related toxicity and mortality.¹⁴ Based on the even further impaired OS of MNGLP patients with confirmed pathogenic variants, it is possible, that at least some of our identified VUS might be pathogenic thus explaining their obvious impact on OS. However, additional functional analyses are needed to further characterize these variants.

Our data support the need to develop specific screening but also treatment protocols for AML patients with underlying MNGLP to both reduce toxicity and improve response to therapy. In addition, genetic counseling of affected families is crucial and specific screening for MNGLP-related genes in HLA-identical or haploidentical family donors for allogeneic HSCT is mandatory.¹⁵

Current guidelines recommend genetic screening for all MNGLP in adults only in the presence of a positive family

history or characteristic physical abnormalities.² Late-onset adult MNGLP patients often present clinically with few uncharacteristic symptoms leading to substantial underdiagnosis.¹⁴ In line with this finding, we were able to retrospectively confirm with the treating hematologist that the two cases with now confirmed pathogenic variants had initially been classified as de novo AML by the treating physician all located at experienced centers. Similar observations were found in a study of young adult patients with MDS¹⁶ where in a relevant subset of patients identified to have underlying SBDS presented with no relevant phenotype beside short body size.¹⁶

Our study was limited by the retrospective nature of this analysis. Gold standard germline samples could not be obtained and had to be substituted by remission samples following induction therapy. Nevertheless, the data showed that samples in remission provide a suitable source to identify variants in MNGLP genes by NGS using the expected VAF cutoff for germline variants (see Supplement).

Given the significant difference of survival in AML patients with and without variants in MNGLP, our study clearly indicates the need of prospective screening for MNGLP in young patients with AML. Age and aberrant karyotype (potentially complemented by TL screening for TBD) might provide simple parameters to trigger genetic screening for inherited MNGLP in addition to the actual recommendations based on mere family history and clinical findings. Along this line, larger trials are needed to weigh the individual parameters and clarify the added value of TL measurement in this setting.

ACKNOWLEDGMENTS

We thank Anne Abels for the excellent technical assistance. This work was supported by a grant of the “Württembergischer Krebspreis 2019” and START Grant (N° 691743, RWTH Aachen University) to FB as well as the “Aachener Krebs-und Leukämiehilfe” and the Deutsche Forschungsgemeinschaft (DFG) through the CRU344 to TB. KK received support from the Task force COVID-19 Forschungsnetzwerk and MWW was supported through BMBF MyPred (01GM1911B) and ERAPERMED GATA2-HuMo (01KU1904).

AUTHOR CONTRIBUTIONS

MK performed the experiments, analyzed and interpreted the data and wrote the manuscript. MC, CR, FS, MK, MB, HS, UP, CM, CDB provided patient samples, clinical data and revised the manuscript. BR, AM, KK, MB, MV performed the experiments and analyzed the data. MWW, SSS performed the experiments and analyzed the data. THB, EJ, FB conceived and planned the study design, interpreted the data, and wrote the manuscript.

DISCLOSURES

THB and FB receive scientific support from RepeatDx, Vancouver. All the other authors have no conflicts of interest to disclose.

REFERENCES

- Arber DA, Orazi A, Hasserjian R, et al. The 2016 revision to the World Health Organization classification of myeloid neoplasms and acute leukemia. *Blood*. 2016;127:2391–2405.
- Nickels EM, Soodalter J, Churpek JE, et al. Recognizing familial myeloid leukemia in adults. *Ther Adv Hematol*. 2013;4:254–269.
- Alter BP, Giri N, Savage SA, et al. Cancer in the National Cancer Institute inherited bone marrow failure syndrome cohort after fifteen years of follow-up. *Haematologica*. 2018;103:30–39.
- Vieri M, Brummendorf TH, Beier F. Treatment of telomeropathies. *Best Pract Res Clin Haematol*. 2021;34:101282.
- Alabbas F, Weitzman S, Grant R, et al. Underlying undiagnosed inherited marrow failure syndromes among children with cancer. *Pediatr Blood Cancer*. 2017;64:302–305.
- Kirschner M, Maurer A, Wlodarski MW, et al. Recurrent somatic mutations are rare in patients with cryptic dyskeratosis congenita. *Leukemia*. 2018;32:1762–1767.
- Richards S, Aziz N, Bale S, et al. Standards and guidelines for the interpretation of sequence variants: a joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology. *Genet Med*. 2015;17:405–24.
- Ventura Ferreira MS, Crysandt M, Ziegler P, et al. Evidence for a pre-existing telomere deficit in non-clonal hematopoietic stem cells in patients with acute myeloid leukemia. *Ann Hematol*. 2017;96:1457–1461.
- Ostergaard P, Simpson MA, Connell FC, et al. Mutations in GATA2 cause primary lymphedema associated with a predisposition to acute myeloid leukemia (Emberger syndrome). *Nat Genet*. 2011;43:929–931.
- Boocock GR, Morrison JA, Popovic M, et al. Mutations in SBDS are associated with Shwachman-Diamond syndrome. *Nat Genet*. 2003;33:97–101.
- Yang F, Long N, Anekpuranang T, et al. Identification and prioritization of myeloid malignancy germline variants in a large cohort of adult patients with AML. *Blood*. 2022;139:1208–1221.
- Wartiovaara-Kautto U, Hirvonen EAM, Pitkänen E, et al. Germline alterations in a consecutive series of acute myeloid leukemia. *Leukemia*. 2018;32:2282–2285.
- Alkhateeb HB, Nanaa A, Viswanatha D, et al. Genetic features and clinical outcomes of patients with isolated and comutated DDX41-mutated myeloid neoplasms. *Blood Adv*. 2022;6:528–532.
- Dokal I, Vulliamy T. Inherited bone marrow failure syndromes. *Haematologica*. 2010;95:1236–1240.
- Ayas M, Nassar A, Hamidieh AA, et al. Reduced intensity conditioning is effective for hematopoietic SCT in dyskeratosis congenita-related BM failure. *Bone Marrow Transplant*. 2013;48:1168–1172.
- Lindsley RC, Saber W, Mar BG, et al. Prognostic mutations in myelodysplastic syndrome after stem-cell transplantation. *N Engl J Med*. 2017;376:536–547.