Somatic mutations of isocitrate dehydrogenases 1 and 2 are prognostic and follow-up markers in patients with acute myeloid leukaemia with normal karyotype

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Background. Mutations in the isocitrate dehydrogenase 1 and 2 (*IDH1* and *IDH2*) genes are frequent molecular lesions in acute myeloid leukaemia with normal karyotype (AML-NK). The effects of *IDH* mutations on clinical features and treatment outcome in AML-NK have been widely investigated, but only a few studies monitored these mutations during follow-up.

Patients and methods. In our study samples from 110 adult de novo AML-NK were studied for the presence of *IDH1* and *IDH2* mutations, their associations with other prognostic markers and disease outcome. We also analyzed the stability of these mutations during the course of the disease in complete remission (CR) and relapse.

Results. *IDH* mutations were found in 25 (23%) patients. *IDH*⁺ patients tend to have lower CR rate compared to *IDH* patients (44% vs 62.2%, p = 0.152), and had slightly lower disease free survival (12 months vs 17 months; p = 0.091). On the other hand, the presence of *IDH* mutations had significant impact on overall survival (2 vs 7 months; p = 0.039). The stability of *IDH* mutations were studied sequentially in 19 *IDH*⁺ patients. All of them lost the mutation in CR, and the same *IDH* mutations were detected in relapsed samples.

Conclusions. Our study shows that the presence of *IDH* mutations confer an adverse effect in AML-NK patients, which in combination with other molecular markers can lead to an improved risk stratification and better treatment. Also, *IDH* mutations are very stable during the course of the disease and can be potentially used as markers for minimal residual disease detection.

 $\textit{Key words: IDH1} \ \textit{mutations; IDH2} \ \textit{mutations; acute myeloid leukaemia; normal karyotype } \\$

Introduction

Patients with acute myeloid leukaemia with normal karyotype (AML-NK) comprise 40-50% of all AML patients.¹ They are characterized by high heterogeneity in terms of clinical features, bio-

logical characteristics and response to treatment. Nevertheless, all of the AML-NK patients are categorized into intermediate risk group. The need for more precise risk stratification of such cases led to the discovery of numerous new molecular markers. Some of them, such as mutations in fms-related

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tyrosine kinase-3 (*FLT3*), nucleophosmin (*NPM1*) and CCAAT/enhancer binding protein alpha (*CEBPA*) genes have made an impact on prognosis of AML-NK patients. Those mutations have been already included in the revised version of World Health Organisation classification of leukaemia.² This new classification implies that all AML-NK patients with mutated *NPM1* without *FLT3*- (internal tandem duplication) *ITD* and mutated *CEBPA* have favourable genotype.

Mardis *et al.* have reported the entire genome sequence of leukemic cells from a single *de novo* AML-NK patient and compared it with the genome sequence from normal skin cells of the same patient.³ After that, from the number of possible somatic mutations, only a handfull of genes were recurrently mutated in multiple AML genomes, including mutations in the genes for isocitrate dehydrogenase 1 (*IDH1*) and isocitrate dehydrogenase 2 (*IDH2*).

The *IDH1* and *IDH2* genes, located at chromosome bands 2q33.3 and 15q26.1 respectively, encode NADPH (reduced nicotinamide adenine dinucleotide phosphate) - dependent isocitrate dehydrogenase 1/2 enzymes, whose main role is to protect cells from oxidative stress.⁴

Heterozygous point mutations in *IDH1* and *IDH2* genes most likely affect the evolutionarily conserved arginine at position R132 in exon 4 of *IDH1* (IDH^{R132}) and either the homologous position R172 (IDH^{R172}) or the second arginine R140 in the IDH2 gene (IDH^{R140}).⁵

IDH1 and *IDH2* mutations occur in approximately 20% of AML-NK cases. ⁶⁻¹¹ Clinical characteristics commonly found in these patients compared to those with wild-type *IDH* are older age, higher platelet counts and concurrent presence of *NPM1* mutations. ^{5,6,8,9,11,18} The relatively high incidence of *IDH* mutations and their association with the most commonly detected mutations in AML patients (*NPM1* mutations) indicates possible mutual interactions in the pathogenesis of the disease. ^{19,22}

Despite the results of numerous studies investigating the effect of the presence of *IDH* mutations on clinical outcome, the prognostic significance of these mutations remains controversial.¹¹ A number of studies showed that the presence of these mutations have no effect in response to therapy and survival^{5,14-16}, while there are others that suggest a negative prognostic effect.^{8-10,17-20} Nevertheless, most studies agree with the fact that *IDH* mutations have adverse prognostic impact in the so called low-risk group of patients (*NPM1**/*FLT3*-*ITD*- AML-NK patients).^{8-10,13,20,21}

Some studies investigated the potential of *IDH* mutations as a follow-up markers. ^{13,16,22-24} *IDH1* and *IDH2* mutations are relatively stable and show direct correlation with disease status. Thus, *IDH* mutations could be useful markers for monitoring disease, including treatment response, minimal residual disease (MRD), and early relapse.

The purpose of our study was to analyze the frequency of mutations in *IDH1*/2 genes and their potential associations with other prognostic markers and outcome in 110 adult *de novo* AML-NK patients. We also analyzed the stability of these mutations during the course of the disease in complete remission (CR) and relapse.

Patients and methods

Patients

From 2009-2014, pre-treatment bone marrow (BM) samples from 110 consecutive consenting patients with de novo AML-NK were analysed at the Clinic for Hematology. This study was approved by the by the Ethics Committee of the Clinical Centre of Serbia, Belgrade, Serbia. Written informed consent was obtained for all patients. Diagnostic procedures comprised cytomorphology, cytogenetics, molecular genetics and immunophenotyping of BM. Morphologic diagnosis was made according to the French-American-British classification.²⁵ Conventional G-band karyotyping was employed for cytogenetic analysis.26 Immunophenotyping by flow cytometry was performed using the direct multicolour immunofluorescent technique applied to whole BM specimens.2 A WBC count ≥ 30x10⁹/L was considered as leukocytosis. Organ dysfunctions, as well as non-disease mortality risk were estimated by the Hematopoietic Cell Transplantation Comorbidity Index (HCT-CI).²⁷ Performance status was assessed using the Eastern Cooperative Oncology Group (ECOG) scale.²⁸ All patients < 60 years of age were treated with standard "3+7" induction chemotherapy, consisting of daunorubicin at a daily dose of 60 mg/m² on days 1–3, in combination with cytarabine at 200 mg/ m² daily as a continuous intravenous infusion for 7 days. Patients > 60 years old were treated with reduced doses in the same regimen. Patients who achieved CR after induction chemotherapy received three cycles of consolidation chemotherapy: cytarabine 3 g/m² per q12h on days 1, 3 and 5 for those younger than 60 years and cytarabine 0.5-1g/m²per q12h on days 1, 3 and 5 for those older than 60 years. Patients aged ≤55 years underwent allogeneic stem cell transplantation (SCT), in total 15 (25.42%) patients. Definitions of CR, overall survival (OS), disease free survival (DFS) and early death (ED) were established by proposed criteria.²⁹

Molecular analyses

BM samples collected at diagnosis, in CR (after induction therapy and after consolidation) and at relapse were analysed. Mononuclear cells were separated by Ficoll density gradient centrifugation and cryopreserved until mutational analyses. Genomic DNA was extracted from the mononuclear cells using a QIAamp Blood Mini Kit (Qiagene, Germany) according to the manufacturer's protocol. DNA fragments spanning exons 4 of the IDH1 and IDH2 genes were amplified by polymerase chain reaction (PCR) as described before.24 PCR reaction products were further subjected to direct sequencing, and the resulting sequences compared to wild-type IDH1 and IDH2 cDNA (GenBank Accession numbers NM 005896.2 and NM 002168.2, respectively). Mutational analyses of FLT3 and NPM1 gene mutations were performed as previously reported.³⁰⁻³² We investigated the impact of IDH mutations on OS in AML-NK patients in relation to three different risk groups defined by FLT3 and NPM1 mutation status (favourable risk-NPM1+/FLT3-ITD-; intermediate-NPM1⁻/FLT3-ITD⁻; unfavorable-FLT3-ITD⁺), according to the recommendation of European Leukaemia Net.1

Statistical analysis

Differences in continuous variables were analysed using the Mann-Whitney U test for distribution between two groups. Frequencies were analysed using the Pearson χ^2 test for 2x2 tables or the Fisher exact test for larger tables. Survival probabilities were estimated by the Kaplan-Meier method, and differences in survival distributions were evaluated using the Log rank test. Patients undergoing allogeneic SCT were censored at the time of transplantation. Multivariate logistic regression model was applied to analyse factors related to the probability of CR failure. Cox's regression model was applied to determine the association of NPM1 mutations with OS and DFS with adjustment for other factors. The statistical analyses were performed using SPSS computer software 15.0 (Chicago, IL, USA). For all analyses, the probability (p) values were 2-tailed and p < 0.05 was considered statistically significant.

TABLE 1. Type of *IDH1* and *IDH2* mutations identified in 110 AML-NK patients

Mutation	Nucleotide change	Amino acid change	No. of patients	
IDH1				
	c.394C>T	R132C	4	
	c.395G>A	R132H	2	
	c.394C>G	R132G	1	
	c.394C>A	R132S	1	
IDH2				
	c.419G>A	R140Q	12	
	c.418C>T	R140W	1	
	c.419G>T	R140L	1	
	c.418C>G	R140G	1	
	c.515G>A	R172K	2	

C = cysteine; G = glycine; H = histidine; K = lysine; L = leucine; S = serine. Q = alutamine

IDH1 and IDH2 nucleotide numbering based upon the NCBI sequence NM_005896.2 and NM_002168.2, respectively.

Results

Frequency of IDH1 and IDH2 mutations in AML-NK patients

Among the 110 AML-NK patients, 25 (23%) harboured missense mutations in IDH genes. Eight (7%) patients had IDH1 mutations, all of them IDH^{R132} . Seventeen (16%) patients had IDH2 mutations: fifteen IDH^{R140} and two IDH^{R172} (Table 1). The wild-type allele was retained in all IDH positive samples, and no patient had both IDH1 and IDH2 mutations. As IDH1 and IDH2 mutations were mutually exclusive and appear to have the same functions, we examined the clinical significance these mutations as a collective group as previously reported. 11

Association of IDH mutations with clinical characteristics and other molecular markers

Pre-treatment clinical characteristics of the patients are summarized in Table 2. Their mean age was 54 years (range 19–78), while 31 (31.8%) patients were \geq 60 years of age. There were 62 (56.4%) men and 48 (43.6%) women. Distribution of IDH^+ patients across FAB groups was uneven, being most frequent in the M2 group - nine (29%) patients, followed by six (27.3%) in the M1 and five (21%) in the M4 group. IDH^+ patients had higher platelet counts (p = 0.024), as well as a higher percentage of pe-

TABLE 2. Clinical characteristics of patients with de novo AML-NK stratified by the presence or absence of IDH mutations

Parameter	Total (n = 110)	IDH+ (n = 25)	IDH ⁻ (n = 85)	p value
Sex				0.617
Male (%)	62 (56.4)	13 (21)	49 (79)	
Female (%)	48 (43.6)	12 (25)	36 (75)	
Age, years, median (range)	53.5(19-78)	50(23-73)	54(19-78)	0.783
ECOG ≥2				0.081
Yes No	45(40.9) 65(59.1)	14(31.1) 11(16.9)	31 (68.9) 54 (83.1)	
HCT-CI ≥3				0.300
Yes No	8(7.3) 102(92.7)	3 (37.5) 22(21.6)	5 (62.5) 80(78.4)	
WBC count, x10°/l (range)	16.8 (0.5-195)	6.9 (0.5-160)	17.4 (0.8-195)	0.373
Haemoglobin median, range	95.5 (6-178)	100 (57-178)	94 (6-140)	0.810
Platelets (x10°/L) median, range	68 (1-420)	109(16-193)	56 (1-420)	0.024
LDH (U/L) median, range	917 (273-7180)	901 (315-5105)	922.5 (273-7180)	0.825
Peripheral blood blast (%)	26 (0-96)	60.5 (0-96)	21 (0-96)	0.031
Bone marrow blasts (%)	71 (23-97)	67 (33-97)	73 (23-97)	0.920
FAB (%)				0.139
MO	10	4 (40)	6 (60)	
M1	22	6 (27.3)	16(72.7)	
M2	31	9 (29)	22 (71)	
M4	24	5 (21)	19 (79)	
M5	22	1 (0.05)	21 (95.5)	
M6	1	0 (0.0)	1 (100.0)	
FLT3-ITD				0.626
present (%)	26(23.6)	5 (19.2)	21 (80.8)	
absent (%)	84(76.4)	20 (23.8)	64 (76.2)	
FLT3-D835				0.428
present (%)	9	3 (33.3)	6 (66.7)	
absent (%)	101	22 (21.8)	79 (78.2)	
NPM1				0.496
present (%)	42(38.2)	11(26.2)	31 (73.8)	
absent (%)	68(61.8)	14(20.6)	54(79.4)	

ECOG = performance status of the Eastern Cooperative Oncology Group; FAB = French-American-British classification; FLT3-ITD = FLT3 internal tandem duplication; HCT-CI = hematopoietic cell transplantation-comorbidity index; IDH = isocitrate dehydrogenase; LDH = lactate dehydrogenase; NPM1 = nucleophosmin; WBC = white blood cell count

ripheral blood (PB) blasts (p = 0.031) compared to IDH^- patients. There were no differences between IDH^+ and IDH^- patients regarding age, sex, WBC count, BM blast percentage, haemoglobin and serum LDH level.

IDH mutations occurred evenly in *NPM1*⁺ and *NPM1*⁻ patients (26.2% vs 20.6%, p = 0.496). Moreover, *IDH* mutations were not associated with *FLT3-ITD* mutations: 19.2% vs 23.8% (p = 0.626).

Response to induction therapy and prognostic relevance of IDH mutations

Out of the 85 *IDH* patients, 51 (62.2%) achieved CR, while 11/25 (44%) of the *IDH*⁺ patients achieved CR. The difference was not statistically significant (p = 0.152). The presence of *IDH* mutations was not associated with ED (*IDH*⁺-36% vs *IDH*- 24.7%; p = 0.310), too. Overall 36/110 (32.7%) participants

exhibited disease relapse, 6 (24%) IDH^+ and 30 (35.3%) IDH^- patients. The impact of IDH mutations on DFS failed to reach statistical significance (IDH^+ - 12 months $vs\ IDH^-$ 17 months; p=0.266). In contrast, OS was significantly impaired in the presence of IDH mutations (IDH^+ -2 months $vs\ IDH^-$ 7 months; p=0.039) (Figure 1).

In the univariate analysis, leukocytosis (p = 0.016) was found to be significantly correlated with a poor rate of CR. The most important factor associated with poor CR rate in the multivariate analysis was leukocytosis (p = 0.015, RR 0.34, 95% CI 0,143-0,809). Univariate analysis showed that significant factors for poor DFS were FLT3-ITD positivity (p = 0.03) and NPM1 positivity (p = 0.032). The most significant risk factor for DFS using the multivariate method was FLT3-ITD positivity (p = 0.030, RR = 2.465, 95% CI 1.089-5.579). Univariate COX proportional regression analysis indicated that the following tested features were significant predictors of poor OS: age \geq 55 years (p = 0.023), leukocytosis (p = 0.001) and IDH positivity (p = 0.039). The multivariate COX proportional regression method pointed to leukocytosis (p = 0.001, RR = 1.768, 95% CI 1.084-2.883) as the most significant predictor of poor OS.

In our study, patients aged 55 years or less received conventional or reduced intensity allogeneic SCT. OS rate in IDH^+ patients not given allogeneic SCT was markedly lower than that in IDH^+ patients who received it (2 vs 15 months; p = 0.006) (Figure 2). Conversely, among patients who did receive allogeneic SCT, the difference in OS rates between those with or without IDH mutations was not significant (p = 0.07).

We found that the presence of IDH^+ had a negative impact on OS in the *intermediate* risk subgroup (5 vs 12 months; p = 0.050) (Figure 3). However, IDH mutations did not affect OS in the *favourable* and *unfavourable* subgroups (1 vs 3 months, p = 0.668; 1 vs 7 months, p = 0.114, respectively).

Sequential studies of IDH mutation

The *IDH* mutational status was serially studied in relapsed samples of *IDH*- patients and in follow-up and/or relapsed samples in *IDH*+ patients. None of the available relapsed samples of *IDH*- patients acquired *IDH* mutations. Among the nineteen *IDH*+ cases who were alive after induction, eleven (44%) achieved CR. Nine of them lost *IDH* mutations after induction therapy but two patients retained it. One of them achieved CR after the first induction therapy. He lost *FLT3-D835* and *NPM1* positive status, but remained *IDH2*+ positive and died

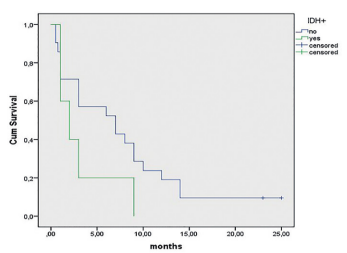


FIGURE 1. Impact of *IDH* mutation on overall survival (p = 0.039 by Kaplan-Meier method).

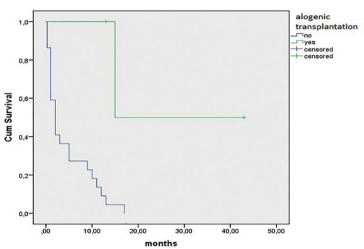


FIGURE 2. Overall survival associated with *IDH* mutations and allogeneic stem cell transplantation in AML-NK patients (p = 0.006 by Kaplan-Meier method).

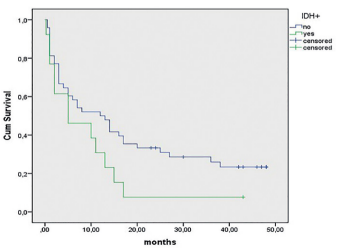


FIGURE 3. Comparison of the overall survival in intermediate group ($NPM1^{-}/FLT3^{-}$) between IDH^{+} and IDH^{-} patients (p = 0.050 by Kaplan–Meier method).

TABLE 3. Results of sequential studies of IDH+ patients

Patient ID	Age/ sex	IDH/FLT3/NPM status on diagnosis	Disease status after induction	IDH/FLT3/ NPM status after induction	Disease status after consolidation therapy	IDH/FLT3/NPM status after consolidation	Relapse	IDH/FLT3/NPM status in relapse
245	53/F	IDH2 ^{R140Q} /wt/wt	CR	wt /wt/wt	CR	/	yes	/
275	50/F	IDH2 ^{R172K} /wt/wt	CR	wt /wt/wt	CR	/	yes	IDH2 ^{R172K} / FLT3-D835/wt
280	61/F	IDH1 ^{R132H} / FLT3-ITD/Type A	RD	/	RD	/	/	/
291	38/F	IDH2R140Q /FLT3-ITD/Type A	RD	/	RD	/	/	/
305	47/M	IDH2R140Qwt/wt	CR	wt /wt/wt	CR	/	yes	/
320	40/M	IDH2R140Wwt/wt	CR	wt /wt/wt	CR	wt /wt/wt	/	/
349	39/M	IDH2R140L/wt/wt	CR	IDH2 ^{R140L} / wt/wt	CR	wt /wt/wt	/	/
378	66/M	IDH1 ^{R132H} /wt/Type A	ED	/	/	/	/	/
380	44/F	IDH1 ^{R132C} /wt/wt	CR	wt /wt/wt	CR	/	yes	IDH1R132C/wt/wt
393	54/M	IDH2R132C/ FLT3-D835/wt	CR	wt /wt/wt	CR	/	yes	/
399	23/F	IDH2R140Q/wt/Type A	ED	/	/	/	/	/
401	69/M	IDH2R140Q/FLT3-ITD/wt	RD	/	/	/	/	/
403	73/F	IDH1 ^{R132C} /wt/wt	RD	/	/	/	/	/
412	46/M	IDH2R140Q/wt/Type A	RD	/	/	/	/	/
418	62/F	IDH1 ^{R132G} /FLT3-ITD/Type A	CR	wt /wt/wt	CR	/	/	/
423	43/M	IDH2R140Q/wt/wt	ED	/	/	/	/	/
426	56/M	IDH2 ^{R172K} /wt/wt	ED	/	/	/	/	/
469	63/M	IDH1 ^{R132C} /FLT3-D835/ Type A	CR	IDH1 ^{R132C} / wt/wt	CR	/	yes	/
487	73/M	IDH2R140Q/wt/wt	RD	IDH2 ^{R140Q} / wt/wt	RD	/	/	/
556	50/M	IDH2R140Q/wt/Type A	ED	/	/	/	/	/
612	30/M	IDH2R140Q/wt/wt	CR	wt /wt/wt	CR	wt /wt/wt	/	/
615	40/F	IDH2R140Q/wt/Type A	CR	wt /wt/wt	CR	wt /wt/wt	/	/
645	43/F	IDH2R140G/ FLT3-ITD/Type A	RD	/	/	/	/	/
672	33/F	IDH1 ^{R132S} /wt/Type A	RD	/	/	/	/	/
680	67/M	IDH2R140Q/FLT3-D835/wt	ED	/	/	/	/	/

CR = complete remission; ED = early death; RF = refractory disease; wt = wild type

in relapse of disease (patient ID 469). The second one (patient ID 349) achieved CR after of induction therapy but retained *IDH* mutation. The mutation was lost in sequential follow-up sample, but patient died during the consolidation therapy in cyto-morphological remission with bone marrow aplasia from the septic shock (Table 3). Patient with refractory disease (patient ID 487) two months after the beginning of therapy remained *IDH2* positive. Two patients, who lost their *IDH* mutation in CR, regained it in relapse. Two of the nine patients

who achieved molecular remission were treated with allogeneic SCT and are still alive. Remaining 7 patients died during therapy and after disease relapse. These results indicate stability of *IDH* mutations during the course of AML.

Discussion

The frequency of *IDH* mutations in patients with AML is 6-19%, but 12-33% in those with AML-

NK. 5,8,17,18,20,33 In our study on adult AML-NK patients, *IDH* mutations were detected in 23% of them. The prevalence of *IDH2* over *IDH1* mutations observed here (15.5% vs 7%) was similar to other published results. 5,6,8,22

Our patients with *IDH* mutations had higher platelet counts and a higher percentage of PB blasts than those without such mutations, which confirms previous findings.^{5,6,8,11} We detected *IDH* mutations most frequently in M2 type cases, followed by M1 (36% and 24%, respectively) and M4 type, which is in according with other results.^{13,15,19}

Examining correlations between *IDH* mutations and other common genetic alterations in AML, such as *NPM1* and *FLT3* mutations, we found a slight but non-significant prevalence of *NPM1*⁺ among *IDH*⁺ patients (*NPM1*⁺- 26.2% vs *NPM1*⁻-20.6%; p = 0.496). This is not in line with previous reports.^{6, 8,9,11,18} The *FLT3* mutations were almost equally distributed between *IDH*⁺ and *IDH*⁻ groups of patients, which is in concordance with other studies.^{5,7,18}

The prognostic impact of *IDH* mutation is controversial. Most studies have shown that both *IDH1* and *IDH2* mutations confer an unfavourable prognosis in AML-NK, *i.e.* a higher risk of disease relapse and shorter OS.^{3,6-8,11,16,17,20} In our study, CR rate was 62.2% in *IDH*⁻ patients, while in *IDH*⁺ patients it was somewhat lower (45.8%), but without statistical significance. A similar finding was reported by Nomdedeu *et al.*¹⁷, where the CR rate of *IDH*⁻ patients was 80% and 63% in *IDH*⁺ (p = 0.086).

We were able to demonstrate that *IDH* mutations act as an adverse prognostic marker of OS in AML-NK patients. That is, patients with *IDH* mutations had significantly worse OS, with a tendency for shorter DFS. This also confirmed earlier findings. ^{6-8,11,16,17,20} Among the *IDH*+ patients, OS rate in those who received allogeneic SCT was significantly higher than that in those not given it. This was also observed by Yamaguchi *et al.*¹¹ and suggests that allogeneic SCT may improve OS in younger patients with *IDH* mutations.

The emergence of new molecular markers in AML-NK has contributed to a better and more precise classification of patients. This group is identified as an intermediate risk group, but because of its heterogeneity in terms of clinical outcome of the disease, more precise allocation is necessary. In addition to the *FLT3* and *NPM1* gene mutations that have already found significance as valuable prognostic factors, the detection of *IDH* mutations has contributed to refined risk classification of AML-NK patients.

When we applied molecular classification based on the presence/absence of *NPM1* and *FLT3* mutations in our cohort of patients, we observed that the presence of *IDH* mutations had an adverse impact on OS in the intermediate risk subgroup (*NPM1*/*FLT3-ITD*-). This finding, already reported by others^{11,16}, argues in favour of testing for *IDH* mutations among AML-NK patients.

The frequent co-occurrence of *IDH* mutations with NPM1 and less often with FLT3 mutations, indicates that such mutations cooperate in the process of leukemogenesis. IDH1 and IDH2 are epigenetic modifier genes involved in DNA methylation and histone modification, and do not completely fit into our current definition of type-I and type-II aberrations, as suggested by the 2-hit theory of cancerogenesis.34,35 Nevertheless, it has been suggested that IDH mutations are an early event in a variety of myeloidneoplasias like myelodisplastic syndrome and myeloproliferative neoplasms (MPN).35,36 In patients with MPN, the acquisition of IDH mutations predicts an increased risk of progression to secondary AML, potentially serving as a marker for early stage transformation.³⁷⁻³⁹ Also, the fact that *IDH* mutations are stable during the course of the disease supports the presumption that their emergence is an early event in malignant transformation.

Even though the prognostic significance of *IDH* mutations has been extensively studied, there are only few reports about their value in MRD monitoring. Thus, Gross et al. 40 and Jeziskova et al. 23 each presented four patients with IDH1 and IDH2 mutations, followed by the investigations of Chou et al.15,21 In all three studies, as in our nine IDH+ patients who were available for sequential analysis, the mutation was lost during CR and reappeared at relapse of the disease as the same type of mutation. Moreover, none of the patients acquired new IDH mutations during relapse. 15,21,23,40 In our study, we registered two IDH+ patients retaining the mutation in CR and during the whole follow-up. Chou et al.22 explained a similar finding through the hypothesis that *IDH* mutations are important in maintaining the leukaemia phenotype through cooperation with other oncogenic mutations, but alone are not sufficient for leukaemogenesis *in vivo*.

IDH1 and *IDH2* mutations have significant potential as MRD markers, assuming that the method applied meets the sensitivity criteria for MRD detection. The usual method for discovering *IDH* mutations is PCR-followed by direct sequencing, with a sensitivity of about 20%.8,14,15,18 Based on this and the fact that *IDH* mutations are heterozygous,

the application of more sensitive methods, such as real-time PCR specific for a given mutation, should be considered for monitoring therapy response and early relapse.

In conclusion, acquired *IDH* mutations are common abnormalities in AML-NK. They confer an adverse effect, especially in patients lacking *NPM1* mutations. In combination with other molecular markers, *IDH* mutational status can lead to an improved risk stratification approach for AML-NK patients. Moreover, *IDH* mutations are stable during the course of the disease and can be potentially used as markers for MRD detection. This could be especially important if specific treatment with *IDH* inhibitors is introduced in everyday practice.

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References

- Döhner H, Estey EH, Amadori S, Appelbaum FR, Büchner T, Burnett AK, et al. Diagnosis and management of acute myeloid leukemia in adults: recommendations from an international expert panel, on behalf of the European Leukemia Net. *Blood* 2010; 115: 454-74.
- Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. ITS Classification of tumours of haematopoietic and lymphoid tissues. Lyon: IARC Press: 2008.
- Mardis ER, Ding L, Dooling DJ, Larson DE, McLellan MD, Chen K, et al. Recurring mutations found by sequencing an acute myeloid leukemia genome. N Engl J Med 2009; 361: 1058-66.
- Ward PS, Patel J, Wise DR, Abdel-Wahab O, Bennett BD, Coller HA, et al. The common feature of leukemia-associated IDH1 and IDH2 mutations is a neomorphic enzyme activity converting a-ketoglutarate to 2- hydrozyglutarate. Cancer Cell 2010: 17: 225-34.
- Abbas S, Lugthart S, Kavelaars FG, Schelen A, Koenders JE, Zeilemaker A, et al. Acquired mutations in the genes encoding *IDH1* and *IDH2* both are recurrent aberrations in acute myeloid leukemia: prevalence and prognostic value. *Blood* 2010; 116: 2122-6.
- Paschka P, Schlenk RF, Gaidzik VI, Habdank M, Krönke J, Bullinger L, et al. IDH1 and IDH2 mutations are frequent genetic alterations in acute myeloid leukemia and confer adverse prognosis in cytogenetically normal acute myeloid leukemia with NPM1 mutation without FLT3 internal tandem duplication. J Clin Oncol 2010; 28: 3636-43.
- Boissel N, Nibourel O, Renneville A, Gardin C, Reman O, Contentin N, et al. Prognostic impact of isocitrate dehydrogenase enzyme isoforms 1 and 2 mutations in acute myeloid leukemia: a study by the Acute Leukemia French Association group. J Clin Oncol 2010; 28: 3717-23.
- Marcucci G, Maharry K, Wu YZ, Radmacher MD, Mrózek K, Margeson D, et al. *IDH1* and *IDH2* gene mutations identify novel molecular subsets within de novo cytogenetically normal acute myeloid leukemia: a Cancer and Leukemia Group B study. *J Clin Oncol* 2010; 28: 2348-55.
- Zhou KG, Jiang LJ, Shang Z, Wang J, Huang L, Zhou JF, et al. Potential application of IDH1 and IDH2 mutations as prognostic indicators in nonpromyelocytic acute myeloid leukemia: a meta-analysis. Leuk Lymphoma 2012; 53: 2423-29.

- Im AP, Sehgal AR, Carroll MP, Smith BD, Tefferi A, Johnson DE, et al. DNMT3A and IDH mutations in acute myeloid leukemia and other myeloid malignancies: associations with prognosis and potential treatment strategies. Leukemia 2014: 28: 1774-83.
- Yamaguchi S, Iwanaga E, Tokunaga K, Nanri T, Shimomura T, Suzushima H, et al. IDH1 and IDH2 mutations confer an adverse effect in patients with acute myeloid leukemia lacking the NPM1 mutation. Eur J Haematol 2014; 92: 471-7.
- Chotirat S, Thongnoppakhun W, Promsuwicha O, Boonthimat C, Auewarakul CU. Molecular alterations of isocitrate dehydrogenase 1 and 2 (*IDH1* and *IDH2*) metabolic genes and additional genetic mutations in newly diagnosed acute myeloid leukemia patients. *J Hematol Oncol* 2012; 5: 5.
- Thol F, Damm F, Wagner K, Göhring G, Schlegelberger B, Hoelzer D, et al. Prognostic impact of *IDH2* mutations in cytogenetically normal acute myeloid leukemia. *Blood* 2010; **116**: 614-6.
- Wagner K, Damm F, Göhring G, Görlich K, Heuser M, Schäfer I, et al. Impact of IDH1 R132 mutations and an IDH1 single nucleotide polymorphism in cytogenetically normal acute myeloid leukemia: SNP rs11554137 is an adverse prognostic factor. J Clin Oncol 2010: 28: 2356-64.
- Chou WC, Hou HA, Chen CY, Tang JL, Yao M, Tsay W, et al. Distinct clinical and biologic characteristics in adult acute myeloid leukemia bearing the isocitrate dehydrogenase 1 mutation. *Blood* 2010; 115: 2749-54.
- Schnittger S, Haferlach C, Ulke M, Alpermann T, Kern W, Haferlach T. IDH1
 mutations are detected in 6.6% of 1414 AML patients and are associated with intermediate risk karyotype and unfavorable prognosis in adults
 younger than 60 years and unmutated NPM1 status. Blood 2010; 116: 548696
- Nomdedéu J, Hoyos M, Carricondo M, Esteve J, Bussaglia E, Estivill C, et al. Adverse impact of *IDH1* and *IDH2* mutations in primary AML: experience of the Spanish CETLAM group. *Leuk Res* 2012; 36: 990-7.
- Patel JP, Gönen M, Figueroa ME, Fernandez H, Sun Z, Racevskis J, et al. Prognostic relevance of integrated genetic profiling in acute myeloid leukemia. N Engl J Med 2012; 366: 1079-89.
- Patel JP, Ravandi F, Ma D, Paladugu A, Barkoh BA, Medeiros LJ, et al. Acute myeloid leukemia with *IDH1* or *IDH2* mutation: frequency and clinicpathologic features. *Am J Clin Pathol* 2011; 135: 35-45.
- Ravandi F, Patel K, Luthra R, Faderl S, Konopleva M, Kadia T, et al. Prognostic significance of alterations in *IDH* enzyme isoforms in patients with AML treated with high-dose cytarabine and idarubicin. *Cancer* 2012; 118: 2665-73.
- Chou WC, Lei WC, Ko BS, Hou HA, Chen CY, Tang JL, et al. The prognostic impact and stability of Isocitrate dehydrogenase 2 mutation in adult patients with acute myeloid leukemia. *Leukemia* 2011; 25: 246-53.
- Chou WC, Peng KY, Lei WC, Ko BS, Tsay W, Kuo CH, et al. Persistence of mutant isocitrate dehydrogenase in patients with acute myeloid leukemia in remission. *Leukemia* 2012; 26: 527-9.
- Jeziskova I, Razga F, Bajerova M, Racil Z, Mayer J, Dvorakova D. IDH2 mutations in patients with acute myeloid leukemia: missense p.R140 mutations are linked to disease status. Leuk Lymphoma 2010; 51: 2285-7.
- Kuzmanovic M, Tosic N, Colovic N, Karan-Djurasevic T, Spasovski V, Radmilovic M, et al. Prognostic impact of NPM1 mutations in Serbian adult patients with acute myeloid leukemia. Acta Haematol 2012; 128: 203-12.
- Bennett JM, Catovsky D, Daniel MT, Flandrin G, Galton DA, Gralnick HR, et al. Proposed revised criteria for the classification of acute myeloid leukemia. A report of the French-American-British Cooperative Group. Ann Intern Med 1985: 103: 620-5
- Shaffer LG, Slovak ML, Campbell LJ. An international system for human cytogenetic nomenclature. Basel: Karger; 2009.
- Sorror ML, Marris MB, Storb R, Baron F, Sandmaier BM, Maloney DG, et al. Hematopoetic cell transplantation (HCT)-specific comorbidity index: a new tool for risk assessment before allogenic HCT. Blood 2005; 106: 2912-9.
- Oken MM, Creech RH, Tormey DC, Horton J, Davis TE, McFadden ET, et al. Toxicity and response criteria of the Eastern Cooperative Oncology Group. Am J Clin Oncol 1982; 5: 649-55.

- 29. Cheson BD, Bennet JM, Kopecky KJ, Buchner T, Willman CL, Estey EH, et al. International Working Group for diagnosis, standardization of response criteria, treatment outcomes, and reporting standards for therapeutic trials in acute myeloid leukemia. Revised recommendations of the international working group for diagnosis, standardization of response criteria, treatment outcome, and reporting standards for therapeutic trials in acute myeloid leukemia. *J Clin Oncol* 2003; 21: 4642-9.
- 30. Kiyoi H, Naoe T, Yokota S, Nakao M, Minami S, Kuriyama K, et al. Internal tandem duplication of *FLT3* associated with leukocytosis in acute promyelocytic leukemia. Leukemia Study Group of the Ministry of Health and Welfare (Kohseisho). *Leukemia* 1997; **11**: 1447-52.
- Yamamoto Y, Kiyoi H, Nakano Y, Suzuki R, Kodera Y, Miyawaki S, et al. Activating mutation of D835 within the activation loop of FLT3 in human hematologic malignancies. Blood 2001; 97: 2434-9.
- Falini B, Mecucci C, Tiacci E, Alcalay M, Rosati R, Pasqualucci L, et al. GIMEMA Acute Leukemia Working Party: cytolasmatic nucleophosmin in acute myelogenous leukemia with normal karyotype. N Engl J Med 2005; 352: 254-66.
- Figueroa ME, Abdel-Wahab O, Lu C, Ward PS, Patel J, Shih A, et al. Leukemic IDH1 and IDH2 mutations result in a hypermethylation phenotype, disrupt TET2 function, and impair hematopoietic differentiation. Cancer Cell 2010; 18: 553-67.
- 34. Kelly LM, Gilliland DG. Genetics of myeloid leukemias. *Annu Rev Genomics Hum Genet* 2002; **3:** 179-98.
- 35. Fröhling S, Scholl C, Gilliland DG, Levine RL. Genetics of myeloid malignancies: pathogenetic and clinical implications. *J Clin Oncol* 2005; **23**: 6285-95.
- Green A, Beer P. Somatic mutations of *IDH1* and *IDH2* in the leukemic transformation of myeloproliferative neoplasms. *N Engl J Med* 2010; 362: 369-70
- Chotirat S, Thongnoppakhun W, Wanachiwanawin W, Auewarakul CU. Acquired somatic mutations of isocitrate dehydrogenases 1 and 2 (*IDH1* and *IDH2*) in preleukemic disorders. *Blood Cells Mol Dis* 2015; 54: 286-91.
- Abdel-Wahab O, Manshouri T, Patel J, Harris K, Yao J, Hedvat C, et al. Genetic analysis of transforming events that convert chronic myeloproliferative neoplasms to leukemias. Cancer Res 2010; 70: 447-52.
- Tefferi A, Jimma T, Sulai NH, Lasho TL, Finke CM, Knudson RA, et al. *IDH* mutations in primary myelofibrosis predict leukemic transformation and shortened survival: clinical evidence for leukemogenic collaboration with *JAKZV*617F. *Leukemia* 2012; 26: 475-80.
- Gross S, Cairns RA, Minden MD, Driggers EM, Bittinger MA, Jang HG, et al. Cancer-associated metabolite 2-hydroxyglutarate accumulates in acute myelogenous leukemia with isocitrate dehydrogenase 1 and 2 mutations. J Exp Med 2010; 207: 339-44.

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Znotrajžilno zdravljenje nepretrganih anevrizem kavernoznega in oftalmičnega segmenta internih karotidnih arterij s sistemom Pipeline

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Izhodišča. Znotraj žilno zdravljenje širokovratnih anevrizem s preusmerjanjem pretoka krvi je relativno nov način zdravljenja. Preusmeritev pretoka dosežemo z žilnimi opornicami, ki imajo znatno gosteje pleteno steno kot klasične opornice. Kri tako pretežno ostaja v lumnu žilne opornice in zmanjšuje hitrost pretoka znotraj same anevrizme. Zastajanje krvi v anevrizmatski vreči vodi do nastanka tromba in posledične izključitve anevrizme iz obtoka. V raziskavi smo želeli oceniti uspešnost uporabe gosto pletenih žilnih opornic Pipeline pri zdravljenju anevrizem s širokim vratom.

Bolniki in metode. V raziskavo smo zajeli 15 bolnikov, ki smo jih zdravili od novembra 2010 do februarja 2014. Pretežni del anevrizem je ležal na karotidni arteriji, intraduralno v oftalmičnem delu žile. Bolnike smo zdravili z gosto pleteno žilno opornico Pipeline (Ev3), ki smo jih postavili preko vratu anevrizme. Uspešnost zdravljenja smo ocenjevali angiografsko in klinično z nevrološkim pregledom.

Rezultati. Kontrolne angiografije, neposredno po postavitvi žilnih opornic, so pokazale upočasnitev pretoka znotraj anevrizmatske vreče. Pri nobenem izmed bolnikov ni prišlo do tehničnih ali kliničnih zapletov med posegom ali po njem. Kontrolne angiografije smo opravljali šest do dvanajst mesecev po posegu. V večini primerov so bile anevrizme v celoti izključene iz obtoka. Nevrološki status bolnikov ob kontrolnih pregledih je bil brez bolezenskih znakov.

Zaključki. Zdravljenje anevrizem z gosto pletenimi žilnimi opornicami Pipeline in preusmeritvijo obtoka je varna in časovno bistveno krajša metoda v primerjavi s standardno metodo z uporabo žilne opornice in elektolizno ločljivih platinastih zank. Nova metoda predstavlja velik napredek pri zdravljenju kompleksnih anevrizem karotidnih arteriji s širokim vratom in bo verjetno nadomestila dosedanji način zdravljenja.

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Somatske mutacije 1 in 2 izocitrat dehidrogenaze so napovedni označevalci bolnikov z akutno mieloično levkemijo z normalnim kariotipom

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Izhodišča. Mutacije genov 1 in 2 izocitrat dehidrogenaze (*IDH1* in *IDH2*) so pogoste molekularne spremembe pri akutni mieloični levkemiji z normalnim kariotipom (AML-NK). Mnogokrat so preučevali vpliv mutacij *IDH* na klinični potek bolezni in izid zdravljenja AML-NK, vendar samo v nekaj raziskavah so sledili spremembe po zdravljenju.

Bolniki in metode. V raziskavo smo vključili 110 odraslih bolnikov z AML-NK, ki so imeli mutacije *IDH1*. Sledili smo povezavo teh mutacij z ostalimi napovednimi označevalci in izidom bolezni. Preučili smo tudi stabilnost teh mutacij ob poteku bolezni, tako pri popolnih odgovorih na zdravljenje kot pri ponovitvah bolezni.

Rezultati. Mutacije IDH smo zaznali pri 25 (23 %) bolnikih. Bolniki IDH^+ so imeli nižjo stopnjo popolnih odgovorov kot bolniki IDH^- (44 % vs 62,2 %; p = 0,152) in nekoliko manjši interval brez ponovitve bolezni (12 mesecev vs 17 mesecev; p = 0,091). Prisotnost mutacij IDH je značilno zmanjšalo celokupno preživetje bolnikov (2 vs 7 mesecev; p = 0,039). Stabilnost mutacij IDH smo sledili pri 19 bolnikih IDH^+ . Izgubo mutacij smo zabeležili pri bolnikih s popolnim odgovorom na zdravljenje, vendar smo enake mutacije zasledili tudi pri bolnikih s ponovitvijo bolezni.

Zaključki. Rezultati raziskave potrjujejo, da so mutacije *IDH* pri bolnikih z AML-NK napovedni dejavnik, ki skupaj z ostalimi molekularnimi označevalci lahko pripomorejo k stratifikaciji teh bolnikov in omogočijo boljši izbor zdravljenja. Mutacije *IDH* so zelo stabilne med zdravljenjem in bi zato lahko bile označevalec za minimalni preostanek bolezni.