Minimal Residual Disease in Acute Myeloid Leukemia

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Abstract

Technological advances in the laboratory have led to substantial improvements in clinical decision-making by the use of pre-treatment prognostic risk stratification factors in acute myeloid leukemia (AML). Unfortunately similar progress has not been made in treatment response criteria, with the definition of “complete remission” in AML largely unchanged for over half a century. Several recent clinical trials have demonstrated that higher sensitivity measurements of residual disease burden during or after treatment can be performed, that results are predictive for clinical outcome and can be used to improve outcomes by guiding additional therapeutic intervention to patients in clinical complete remission but at increased relapse risk. We review here these recent trials, the characteristics and challenges of the modalities currently used to detect minimal residual disease (MRD), and outline opportunities to both refine detection and better clinically utilize MRD measurements. MRD measurement is already the standard of care in other myeloid malignancies such as chronic myelogenous leukemia (CML) and acute promyelocytic leukemia (APL). It is our belief that response criteria for non-APL AML should be updated to include assessment for molecular complete remission (mCR) and that recommendations for post-consolidation surveillance should include regular monitoring for molecular relapse as a standard of care.

1) Introduction

Our understanding of heterogeneity and the underlying disease biology of the acute myeloid leukemia (AML) has been greatly deepened by the recent use of modern laboratory advances1–4 which have in turn allowed more refined pre-treatment risk stratification using molecular prognostic5, 6 and predictive5 biomarkers. Unfortunately this molecular revolution has been incompletely translated. Just as the most commonly used induction chemotherapy treatment for adult non-APL AML has not changed substantially in the past forty years7, 8 the treatment response criteria for a “complete remission” (CR) has not dramatically changed in almost sixty years9, 10 (Table 1).

It has been recognized since at least 1969 that a clinical “complete remission” is an early, important, necessary but insufficient step on the path to long-term disease control in acute
myeloid leukemia (AML). While a patient at first presentation with high blast count AML may have up to $10^{12}$ leukemic cells (equivalent to several pounds of tumor), that same patient after induction chemotherapy treatment judged to be in an initial “complete remission” may have anywhere as few as none and as many as $10^{10}$ leukemic cells remaining (a number of cells equivalent to a two cubic centimeter tumor mass in a solid tumor setting). It is clear that due to the low sensitivity criteria used that some “complete remissions” are more complete than others (Figure 1). Those in this highly diverse state of “complete remission” currently have clinical decisions regarding subsequent treatment (to establish if the risks associated with stem cell transplantation (SCT) as consolidation therapy are outweighed by a potential benefit from reduction in relapse risk) made by stratification solely on the basis of pre-treatment risk factors rather than any direct quantification of remaining leukemic disease burden.

Many patients diagnosed with AML are not eligible, due to age or comorbidity, for intensive chemotherapy. In those who can tolerate such treatment however initial CR rates of greater than 65% in newly diagnosed poor risk leukemias and up to 92–99% in those with core binding factor (CBF) AML are now achievable with optimal induction therapy which has shifted the focus to the prediction and prevention of relapse.

The current “standard of care” pre-treatment risk factor stratification assigns patients in an initial complete remission to a risk of relapse based on the average clinical outcomes of large historical populations who had similar pre-treatment cytogenetic and molecular characteristics in their leukemia at presentation and does not include any patient-personalized assessment of induction treatment efficacy by high-sensitive detection of residual leukemia burden at the time that decisions regarding most appropriate consolidative treatment must be made.

This ability to measure minimal residual disease (MRD) in AML was not available in 1956 when criteria for the evaluation of response to treatment in acute leukemia were proposed and so understandably thresholds were set based on the technology available at that time. MRD has however been measurable in AML patients in a clinical CR via quantitative reverse transcriptase polymerase chain reaction (RQ-PCR) or flow cytometry (FC) for over twenty years now. Technologies used to detect MRD in acute leukemia have included traditional expert microscopy for morphology, fluorescence in situ hybridization (FISH), cytogenetics, flow cytometry (FC), gene expression analysis using RQ-PCR (GE-PCR), PCR for translocations, rearranged or mutant sequence (Mut-PCR), DNA sequencing and in the setting of monitoring post stem cell transplantation (SCT) analysis of chimerism by PCR or FISH (Figure 1, Table 2, reviewed in). There is already excellent evidence for clinical decision making as a standard of care based on multi-parameter flow cytometry (FC) in acute lymphoblastic leukemia (ALL) and reverse transcriptase-polymerase chain reaction analysis (RT-PCR) in Acute Promyelocytic Leukemia (APL) so the remainder of this review will therefore focus solely on the use and opportunities for use of MRD in non-APL AML.

In this article we compare the modalities of MRD detection currently in use in AML by reviewing twelve recently reported clinical trials or series (Table 3) for what they teach us
regarding the optimal technology, indication, timing and predictive and prognostic utility of such MRD measurements. We conclude by discussing clinical scenarios where we believe MRD measurement in AML could have immediate utility.

2) MRD Detection by Gene Expression Analysis

The development of quantitative real time polymerase chain reaction (RQ-PCR) technology has provided many logistical and technological advantages compared to other MRD detection methodologies (Table 2, 20, 29, 30). The adaptation of this highly sensitive and specific technology to the quantification of disease burden in chronic myeloid leukemia was responsible for fundamental contributions to drug development, response criteria and treatment guidelines and detection and analysis of treatment failures in that disease. Unfortunately no single rearranged or mutated target sequence analogous to the \textit{bcr-} abl fusion transcript is found in all cases of AML, motivating the search for genes expressed at higher levels in AML than normal tissues that could serve as a target for RQ-PCR. While a variety of such genes (eg: leukemia associated antigens) are known and could be potentially used for this purpose, much of the work has focused on the use of \textit{WT1}.

\textit{WT1} is a zinc finger transcription factor with an essential role in normal development with expression in the adult restricted to the renal glomerular podocytes and a subset of hematopoietic precursors. It is overexpressed at the mRNA level in 80–90% of AML cases at diagnosis in both peripheral blood (PB) and bone marrow aspirate (BM), which has lead to its adoption as a standardized method for MRD detection using RQ-PCR. This European LeukemiaNet study found the magnitude of \textit{WT1} log reduction after induction chemotherapy to be an independent predictor of relapse (<2 log reduction was associated with a 75% risk of relapse at 5 years compared to only 40% risk in those with a ≥2 log reduction), unfortunately only 46% of 238 patient peripheral blood pretreatment samples examined had \textit{WT1} sufficiently overexpressed to allow the detection of a reduction of that magnitude. PB appeared preferable to BM for subsequent monitoring as higher levels of \textit{WT1} mRNA expression in healthy donor BM (median 19.8 \textit{WT1} copies/10$^4$ \textit{ABL}, range: 0–213) compared to healthy donor PB (median 0.01 \textit{WT1} copies/10$^4$ \textit{ABL}, range: 0.01–47.6) limited the working dynamic range for detection. Patients with \textit{WT1} levels above the upper limit of normal (250 copies/10$^4$ \textit{ABL} in BM or 50 copies/10$^4$ \textit{ABL} in PB) after consolidation therapy were found at a significantly increased risk of relapse (67% vs. 42% at 5 years).

Within the past year there have been some direct “head-to-head” comparisons of \textit{WT1} RQ-PCR assessment with other MRD modalities. Kwon et al., described 21 AML patients followed after allogeneic SCT using MRD monitoring with \textit{WT1} RQ-PCR gene expression analysis, chimerism and flow cytometry. RQ-PCR was analyzed using ddCT method using GUS reference gene and K562 calibrator sample with >0.025% and >0.55% considered positive results after two consecutive measures in PB and BM respectively. Four-color flow cytometry was performed, using least 10$^6$ BM cells, with a leukemic aberrant phenotype of >0.1% considered positive. All of nine relapses (range 50–560 days after SCT) were detectable using \textit{WT1} monitoring, in eight cases this “molecular relapse” preceded clinical (ie: hematological) relapse by a median of 137 days (range: 52–462). Three of these patients

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had continuously elevated WT1 levels (ie: molecularly refractory) while six were initially negative after SCT followed by a molecular relapse and hematological relapse. Eight of these nine patients had molecular relapse detectable from peripheral blood sampling, including all three patients with extra-medullary relapse. In no case was relapse detected by flow cytometry or mixed chimerism prior to WT1 positivity (although it should be noted that fewer timepoints were examined using these methodologies). A tenth patient also had elevated WT1 levels after SCT but had not relapsed by the time of publication. All eleven patients without WT1 overexpression after SCT remained relapse free.

Miyazaki et al. recently compared the use of FC with quantitative PCR for WT1 and/or leukemia-specific fusion transcripts in 41 acute leukemia patients after allo-SCT. In their analysis of 156 samples they demonstrated good concordance (71.8%) between FC and WT1 PCR based MRD detection, with most of the discordance from samples that were PCR positive for WT1 but negative by FC. Unfortunately this report included a mixture of patients with AML (n=31) and ALL (n=10) making additional interpretation challenging.

Rossi et al. reported in 2012 the use of six color flow cytometry with >0.1% leukemia-associated immunophenotypes (LAIPs) in BM considered positive MRD compared to WT-1 gene expression profiling using RQ-PCR. In this single hospital series, of 47 consecutive AML patients achieving a CR after induction chemotherapy, 23 were eligible for this study as they did not have leukemia-specific rearranged or mutated genetic sequence suitable for MRD assessment. This study concluded that FC and WT-1 RQ-PCR “showed comparable capacity in terms of technical performance and clinical significance to identify high risk patients who eventually relapsed”, a surprising finding given that testing was not restricted to those ~50% of patients with WT-1 mRNA elevated to a level where a two log variation could be detected (the mean expression of 3320 copies WT1/10^4 ABL in diagnosis samples would allow on average only slightly more than 1 log dynamic range dramatically limiting the sensitivity). RQ-PCR testing was performed on bone marrow, and the threshold for positive MRD (90 copies/10^4 ABL) was set almost two thirds lower than reported in the prior standardization of this assay. Given these factors it is possible that this study compared FC with a conservative estimate of the optimal performance characteristics of the WT-1 assay.

3) MRD in core binding factor AML

The core binding factor (CBF) AMLs, defined as those carrying either the t(8,21) chromosomal translocation (RUNX1-RUNX1T1 fusion gene) or Inv(16)/t(16;16) rearrangement (CBFB-MYH11 fusion gene), like APL AML, represent ideal AML subtypes for RQ-PCR based MRD monitoring due to presence of a defined uniquely leukemia-specific stable molecular target. The CBF AMLs represent up to approximately 20% of all AMLs in adults and are considered to be “better-risk” or “favorable risk” status based on cytogenetics and molecular stratification. Additional prognostic information in these subsets would be easily actionable; chemotherapy is often preferred for consolidation so treatment escalation to stem cell transplantation would be an option for those at the highest risk of relapse. Additionally, compared to other AML subtypes
effective salvage therapy is available for Inv(16) AML making post-treatment monitoring for early detection of relapse particularly important\textsuperscript{55}.

The AML02 trial\textsuperscript{56} was a multicenter trial that treated 216 children for AML using risk directed therapy based on presentation leukemia genetic factors and FC based MRD which achieved impressively high rates of CR (94\% after induction 2), 3 year event free survival (EFS) (63\%) and 3 year overall survival (OS) (71\%). FC-based MRD levels after induction therapy were important predictors of relapse, with FC MRD >1\% after induction 1 identified in multivariate analysis as being an independent adverse prognostic factor for both EFS and OS. A subsequent report in 2012\textsuperscript{57} retrospectively analyzed 508 samples from 77 of these patients who had leukemia associated rearranged sequence (RUNX1-RUNX1T1, CBFB-MYH11 or MLL fusion transcripts) suitable for detection by PCR. There was good correlation between negative PCR results and FC, of 311 samples classified negative by PCR 308 were negative by FC. Discrepancy was observed between positive PCR results and FC, only 27 of 197 positive PCR results were confirmed by FC, this however may be an artifact of the technology used to process earlier samples, when modern RQ-PCR technology was used there was excellent correlation between PCR and FC results in 138 of 139 samples tested.

The British MRC15 trial also reported in 2012 the results of prospective assessment of MRD using RQ-PCR for RUNX1-RUNX1T1 or CBFB-MYH11 transcripts in 278 patients aged 15–70 treated for CBF AML\textsuperscript{16}. MRD was informative not only in detection of molecular relapse preceding hematological relapse in serial monitoring during remission, but also the result of a single MRD assessment after completion of induction therapy while in complete remission was itself prognostic for subsequent relapse risk in a multivariate analysis that included age, WBC, secondary AML, performance status and sex. After completion of induction chemotherapy to CR those 46\% of patients with t(8,21) who had achieved a 3 log reduction in BM MRD had a cumulative incidence of relapse (CIR) of only 4\% (1 relapse in 28 patients) compared to a CIR of 32\% in the patients who did not achieve this level of initial leukemic tumor burden reduction. Absolute levels of MRD were also significant with a threshold of <100 copies in BM (47\% of patients) identifying a cohort with a CIR of just 7\% and the threshold of 1000 copies in PB was able to discriminate patients after induction into low (78\% of patients, CIR 15\%) and high (22\% of patients, CIR 50\%) risk groups.

MRD assessment while in CR after induction therapy for inv(16) AML also allowed patients to be stratified based on PB copy number into low (<10 copies, 51\% of patients, CIR 21\%), intermediate (10–500 copies, CIR of 56\%) and high (CIR of 100\% for those with >500 copies) risk groups for subsequent relapse. Subsequent serial monitoring during remission after completion of all therapy for t(8,21) detection of >100 copies of RUNX1-RUNX1T1 transcript in PB or >500 copies in BM (ie: molecular relapse) was associated with a hematological relapse rate of 100\% compared to only 7\% in those below this threshold. Similar thresholds could be established as significantly prognostic of relapse risk for Inv(16) patients in CR after completion of therapy by MRD monitoring in BM (>50 copies, 32\% of patients, 100\% hematological relapse rate compared to a CIR of 10\% in those with <50 copies) and PB (>10 copies, 28\% patients, 97\% hematological relapse risk compared to a CIR of 7\% in those with <10).
Finally, the French CBF-2006 trial reported in early 2013 the results of treatment of 198 adults aged 18–60 with newly diagnosed CBF AML using prospective evaluation of pretreatment leukemia molecular factors (FLT3, KIT and RAS mutations) with MRD assessment for relapse risk-stratification. MRD levels were determined from BM using RQ-PCR prior to three monthly consolidation cycles of cytarabine of 3000 mg/m2/12hr on day 1, 3 and 5 followed by GCSF from day 8 to neutrophil recovery. Patients were randomized to one of two induction regimens resulting the initial post-induction conventional (hematological) CR rates of 98–100%. MRD assessment however could distinguish between the efficacy of these induction regimens, demonstrating differences in the depth of remission and the residual disease burden after induction (0.10% vs. 0.26% fusion transcript ratios) in patients in CR, a difference which appeared to persist after subsequent consolidation cycles. Patients in CR who had not achieved a 3 log reduction in MRD after the first cycle of consolidation therapy became eligible on this protocol for stem cell transplantation, that level of reduction was not reached in 24% and 35% of patients on the two arms. Univariate analysis of prognostic factors of relapse risk identified higher white blood cell count (WBC), KIT mutations, FLT3 mutations, less than a 3 log reduction in MRD after first cycle of consolidation (MRD2) and an absolute MRD2 level of >0.1% as associated factors. Multivariate analysis of WBC, KIT and FLT3 mutations, and MRD2 response showed that MRD2 response (by 3 log reduction or absolute level) was the sole factor influencing specific hazard of relapse both in the whole patient cohort, and also in the individual t(8,21) and Inv(16) subsets. This suggestion that when prognosticating relapse risk the reality of disease burden after treatment (MRD) may trump molecular markers of disease biology assessed prior treatment is provocative, and counter to current clinical practice.

4) PCR for mutated or rearranged sequence for MRD in non-CBF AML

As seen with CML, APL, and CBF AML, rearranged or mutated genetic sequence targets allow for high sensitivity RQ-PCR based detection of leukemic burden. While no universal target exists for AML several gene mutations are common. Nucleophosmin (NPM1) gene mutations are seen in 25–30% of cases of AML and appears to be a stable marker for MRD monitoring. The German-Austrian AML study group recently reported on the prospective use of NPM1mut MRD in 245 adult (age 19–61) AML patients. The degree of disease burden as measured by NPM1mut RQ-PCR MRD was significantly associated with prognosis for patients in a complete remission after each cycle of treatment. For example, MRD negativity after completion of induction therapy was associated with a 4 year CIR of only 6.5% (achieved by 26 patients) compared to a CIR rate of 53% in the 111 patients in morphological CR but not achieving this molecular CR (mCR) milestone. Additionally, 4 year CIR was 15.7% (OS: 80%) in the 62 CR patients who were MRDneg are the completion of therapy compared to a CIR of 66.5% (OS: 44%) in those in CR but MRDpos. While BM MRD assessment was preferable during therapy, PB MRD showed good concordance (88%) with BM in post treatment monitoring, when exceeding a threshold of 200 copies NPM1mut/10⁴ ABL was 100% predictive of relapse. Of note, only 9% of evaluable relapses had no or minimally detectable NPM1mut at the time of relapse, confirming the relative stability of this target. Similar results were shown by Miglino et al.
where achievement of a NPM\textsuperscript{mut} molecular CR after induction was associated with a statistically significant lower relapse risk than those who remained with detectable disease, and a statistically significant higher overall survival at 36 months (64.3\% vs. 11.9\%)\textsuperscript{61}. NPM1\textsuperscript{mut} molecular relapse was invariably associated with eventual hematological relapse\textsuperscript{61}. Very recent data from a large 174 patient dataset has determined that specific NPM1\textsuperscript{mut} MRD threshold levels can be determined in CR1, during surveillance after chemotherapy and after allo-SCT that are significantly associated with poor overall and disease free survival. Importantly, this work also showed that sampling from peripheral blood can take the place of bone marrow testing for this MRD assay\textsuperscript{62}.

Despite evidence that the approximately 50\% of AMLs with sufficient WT1 overexpression for MRD monitoring includes many cases of NPM1\textsuperscript{mut}, Inv(16) or adverse cytogenetic AML\textsuperscript{48}, it is nevertheless likely that a validated and stable molecular target for RQ-PCR MRD monitoring can be found for the majority (perhaps 65–75\%) of patients diagnosed with AML. Additional stable molecular markers with potential for development for MRD monitoring include DNMT3A\textsuperscript{63} and CEBPA\textsuperscript{64}. Despite initial concerns that tracking FLT3-ITD mutation would not be informative due to concerns regarding stability\textsuperscript{65} using more sophisticated techniques\textsuperscript{66} this mutation has been shown to be highly predictive for relapse risk\textsuperscript{67, 68}. It is likely therefore that clinical laboratories will have to offer, or contract with a reference laboratory for, a portfolio of RQ-PCR MRD assays, which will increase as next generation sequencing (NGS) technology increases the number of viable molecular targets\textsuperscript{3}. Once whole genome and/or exome sequencing of tumor and germline becomes possible routinely at cancer diagnosis\textsuperscript{69} this will facilitate identification of not only patient-specific MRD markers\textsuperscript{70}, but also allow molecular diagnosis\textsuperscript{71}, subsequent retrospective discovery of prognostic factors\textsuperscript{5}, and triage to the most appropriate trials of targeted agents\textsuperscript{72}. Finally, the ability to identify stable molecular targets within the leukemic stem cell population\textsuperscript{73–75} for MRD monitoring and therapeutic targeting is of obvious great interest.

5) Flow Cytometry for MRD in AML

Flow cytometry is, when used by appropriate experts, an excellent method for determining MRD in AML with high sensitivity and specificity\textsuperscript{22, 56, 76–86}. Some of the advantages and logistical challenges associated with this method are outlined in Table 2. Modern flow cytometry based MRD detection is most commonly based on the definition aberrant cell surface marker expression in AML cells as a leukemia associated phenotype (LAP or leukemia associated immunophenotype, LAIP) that is not, or only infrequently, seen in normal or recovering bone marrow or blood\textsuperscript{79, 87}. These can include asynchronous expression of early and late antigens, cross lineage infidelity (eg: expression of lymphoid markers on myeloid cells) and antigen under or over expression\textsuperscript{79, 81}. Such LAIPs can be personalized to each patient at presentation, with the caveat that AML is a clonal disease and relapsed disease may have a different immunophenotype\textsuperscript{80, 88–91}.

The main advantage of flow cytometry in MRD monitoring is its near universal applicability; one recent study\textsuperscript{92} found that 94\% of patients had a suitable LAIP for use, with the range reported in the literature when at least three color FC has been used ranging between 60–100\%\textsuperscript{79}. We avoid in this review the ungainly terminology of “immunological

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relapse” and “immunological complete remission” when describing the reappearance or absence of MRD as detected by flow cytometry, the levels of MRD necessary for the molecular response criteria described by Hokland and Ommen \(^{23}\) could, however, in principle, also be detected by modern flow cytometry. It is unclear however if quantitative MRD results can be compared across multiple centers given the importance of human and technical factors \(^{24}\) in interpretation, indeed even in the context of a recent clinical trial with identical standard operating procedures, reagents and instruments used variation was seen between four reference laboratories \(^{81}\).

Given their extremely successful experience of using FC for ALL MRD \(^{25}\) those treating pediatric patients have been leaders in using this technology in AML, including the AML02 study already discussed \(^{56}\) where subsequent analysis determined that flow cytometry based MRD measurement after first and second induction was statistically associated with five year event-free and overall survival \(^{57}\).

A cohort of 219 AML patients under the age of 21 treated on the Childrens Oncology Group (COG) trial AAML03P1 was recently analyzed \(^{93}\) for MRD using four color FC using a non-LAIP based “difference from normal” approach \(^{94}\). In 188 patients with initial morphological CR after the first induction phase the presence of FC defined MRD (seen in 25% of patients) was highly correlated with relapse (3yr relapse risk 60% vs. 29% in MRD\(^{neg}\) patients). Cytogenetic and molecular risk factors were associated with risk of MRD while in CR after first induction with 11%, 29% and 50% of those with favorable, intermediate and high-risk cytogenetics being positive for MRD at this timepoint and 25% of those with FLT3-ITD being MRD\(^{pos}\) compared to 0% of those with NPM1 mutations. A multivariate analysis that included molecular risk group stratification and presentation WBC as variables found that MRD positivity after first induction was an independent predictor of relapse free survival (RFS) for those patients in morphological CR. Interestingly of the 27 patients who did not achieve a morphological CR (>5% blasts) after first induction therapy, seven were found to be negative for residual disease by FC and clinically remain long-term survivors.

Flow cytometry for MRD in adult AML patients can also be prognostic. A retrospective analysis of 123 adults enrolled in EORTC/GIMEMA protocols in the period 1998–2008 in a single center, showed that FC-based MRD at the end of consolidation could blur distinctions between pre-treatment cytogenetic and molecular risk groups \(^{95}\). For example, after consolidation, patients in CR from good risk karyotype AML but MRD\(^{pos}\) had a 4 year RFS of only 15% compared to those in CR and MRD\(^{neg}\) from intermediate karyotype AML who had a 4yr RFS of 63%. The use of post-consolidation FC-based MRD status combined with pre-treatment cytogenetics and FLT3 mutational status could segregate patients into groups with low and high risk for subsequent relapse. Further analysis focused on the effect of autologous (auto-SCT) or allogeneic (allo-SCT) stem cell transplantation in the 79 patients who received one of these therapies as post-consolidation therapy. There was no statistical difference in outcome in low risk patients (but 5 non-relapse, transplant related, deaths in the allo-SCT group) but superior outcomes after allo-SCT for patients in the high
risk category (RFS 47% vs. 13% auto-SCT) suggesting MRD-directed therapy escalation can improve outcomes.

Within any individual patient AML is not a single homogenous disease at the level of genetics and biology. Importantly Gerber et. al., recently identified using FC a population of cells that while rare in leukemia at diagnosis, appeared significantly overrepresented in the MRD fraction in patients in a morphological CR after treatment. These cells were capable of producing complete leukemic engraftment in an immunodeficient mouse model of stem cell function. The presence of such cells in patients after treatment was highly correlated with subsequent clinical relapse. This important finding is now the basis of an ongoing clinical trial at Johns Hopkins Hospital to determine the 2year RFS of AML patients with these “leukemic stem cells” (LSC) at first CR who receive either chemotherapy or stem cell transplantation (NCT01588951). The tracking of LSC MRD represents another unique advantage of flow cytometry until such a time when unique molecular markers of such cancer stem cells can be identified; this is an area of much interest.

6) Use of MRD in the setting of Allogeneic Transplantation

Allogeneic stem cell transplantation remains the first and most effective immunotherapy used for hematological malignancies and the possibility of using MRD to enable both appropriate triage of patients pre-transplantation based on their individual risk of relapse and to follow patients and treat prior to frank clinical/hematological relapse is of obvious interest. In addition to those studies already discussed there are several other reports that directly address the question of the role of MRD testing in the peri-transplant setting. Walter et. al., showed, using a “difference from normal” FC based MRD approach, in patients in conventional CR prior to allo-SCT that 2yr relapse rate (64.9% vs. 17.6%) and overall survival (30.2% vs. 76.6) were markedly different in the group positive and negative for MRD respectively. This increased mortality risk in patients MRDpos before allo-SCT remained statistically significant even when adjusting for other factors.

This worse outcome in patients who are in conventional CR but are MRDpos prior to allo-SCT has also been seen when using cytogenetic and WT-1 PCR MRD testing. For example, Pozzi et. al., just reported on the use of WT1 PCR to predict relapse risk in 122 AML patients the allo-SCT setting. They found, using a threshold of 100 WT1 copies/10^4 copies of ABL, that MRD testing using WT1 levels pre-transplant (69 MRDneg patients had relapse rate of 26% vs. relapse rate 53% in 53 patients who were MRDpos), and especially post-transplant (only 16% of 55 MRDneg patients relapsed, compared to 55% of 67 MRDpos patients) was highly predictive of relapse risk. Combining information from pre-and post- transplant timepoints appeared to be especially informative.

7) Can MRD guide therapy to improve outcomes?

Beyond providing patients with prognostic information, detection of residual disease is only truly useful if therapeutic intervention is available that can change the course of the disease. Evidence for effective MRD-based risk-directed therapy has already been discussed; in AML02 levels of MRD during induction were used to determine timing and need for...
escalation of induction therapy and then a combination of initial risk classification and MRD based assessment of induction response was used to assign appropriate consolidation therapy. This MRD-directed approach resulted in superior outcomes compared to prior trials with 3 year EFS (63%) and OS (71%).

Multiple reports of treatment of molecular relapse prior to clinical/hematological relapse have now shown that MRD-directed therapy can lead to improved clinical outcomes. CD34+ donor chimerism analysis is a specific method for monitoring for MRD after allo-SCT with a chimerism level falling below 80% after SCT having been shown to be a harbinger of hematological relapse in a median of 61 days. The RELAZA trial demonstrated that 4 cycles of azacitidine therapy for MDS/AML patients with CD34+ chimerism falling below this level could delay hematological relapse to a median of 231 days and lead to continuous RFS in 20% of patients without any further treatment. Analogously, it has been shown that rising NPM1$^{\text{mut}}$ MRD levels are predictive of upcoming hematological relapse in a median time of eight weeks. Sockel et al., treated ten AML patients in CR but with increasing NPM1$^{\text{mut}}$ MRD levels with azacitidine therapy with a resulting molecular response (at least 1 log reduction in MRD levels) in 7 of these patients and 7 patients remaining in a complete hematological response after a median follow-up of ten months.

In the Pozzi study already discussed retrospective analysis suggested that patients with rising WT1 MRD who received immunotherapy intervention in the form of donor lymphocyte infusions (DLI) had a better 5 year overall survival (44%) than those with rising WT1 MRD levels who did not receive DLI. This non-randomized study however has several confounding factors however that make interpretation challenging, exacerbated by the fact that time to hematological relapse was not different between the two groups. There is strong evidence however that DLI can be an efficacious intervention in the setting of MRD relapse. Yan et al. analyzed data from 814 patients aged 2–59 receiving SCT (for AML, ALL or MDS-RAEB) from a single center in the period 2006–2010. The 105 patients who were MRD$^{\text{pos}}$ by FC or WT1 RQ-PCR after SCT were treated with either IL-2 and/or modified DLI. Presence of MRD after SCT was associated with greater 3yr CIR (46% vs. 18.1%). Significant differences at 3years were seen within the MRD$^{\text{pos}}$ group based on the modality used to treat their molecular relapse; CIR of 64.4% (OS of 28.1%) in those treated with IL-2 alone compared to CIR of only 27.8% (OS of 58.3%) in those treated with DLI. In fact, CIR and disease free survival was not statistically different between the MRD$^{\text{neg}}$ patients and those MRD$^{\text{pos}}$ patients treated with DLI. Unfortunately responses for MDS/AML patients were not reported separately from those in ALL patients.

The same group has just reported their experience of MRD-directed therapy for patients with t(8,21) AML. MRD determined by RUNX1/RUNX1T1 transcript levels by RQ-PCR was shown to be predictive for relapse when measured after the second cycle of consolidation. A non-randomized comparison of those in a high relapse risk group (defined by being in a morphological CR but MRD$^{\text{pos}}$ after 2nd consolidation) who received allo-SCT as mandated by risk-directed protocol (n=40) compared with those who did not due to patient preference or lack of appropriate donor (n=29) showed a substantial benefit to this escalation of therapy (CIR 22.1% vs. 78.9%, p<.0001). This benefit to allo-SCT was not seen however in the low risk (CR, MRD$^{\text{neg}}$ after 2nd consolidation) group. These observations need confirmation in a
randomized trial but add to the increasing body of evidence that not only is MRD prognostic, but that therapy based on MRD may improve clinical outcomes.

8) Conclusion – how MRD can help now

Despite the challenges in standardizing definitions of assay and AML specific thresholds of MRD and the lack of one universal modality or target, there is now ample indisputable evidence that the MRD monitoring can improve relapse risk stratification and detection in AML. The use of MRD to guide treatment is already an accepted standard of care in acute lymphoblastic leukemia, CML, and APL and based on recent studies discussed here will also soon be in CBF AML.

In the recent CBF AML MRD trials the average kinetics of relapse appeared to be approximately 0.5–0.6 log per month, MRD was checked 3–4 monthly in peripheral blood in the first two year period when most relapses occurred and the median lead time between molecular relapse and hematological relapse was 3 to 4.5 months. While disease biology may influence the kinetics of relapse and the sensitivity of the MRD detection modality used will influence the lead-time between molecular and hematological relapse it now seems clear that regular MRD monitoring could be easily integrated into the current recommendations for post-consolidation surveillance of a peripheral blood draw for complete blood count with platelets every 1–3 months for the first two years and that doing so would provide a therapeutic window for intervention before frank clinical relapse.

The search for optimal MRD assessment in other AML types will be an iterative process and the understandable quest for the perfect should not become the enemy of the introduction of the better; even low sensitivity MRD to a modest 10⁻³ or 10⁻⁴ detection threshold (see figure 1) may provide more actionable information than traditional hematological CR. There are a number of specific clinical scenarios were the adoption of routine standard of care MRD monitoring could make an immediate impact (Table 4).

MRD thresholds need not be infallibly associated with evitable hematological relapse to be clinically useful, the concept of maintenance therapy for prevention of relapse in those at increased risk is already under investigation in myeloma, follicular lymphoma and lung cancer and as less toxic therapies with efficacy in leukemia become available identification of patients at the greatest risk of relapse who may benefit from such additional therapy in the post-conventional treatment setting would be particularly useful. Similarly, monitoring for the early detection of the development of subclinical disease would also have utility for those therapies including immunotherapies where optimal effects might be observed in lower disease burden states.

Standardized mCR criteria will likely also aid drug development. Hematological CR is a blunt tool to evaluate differences between therapies and MRD measurement offers greater dynamic range to detect changes in leukemia burden and monitor for treatment effect on leukemic subpopulations that may be associated with relapse. The ability to dose escalate or de-escalate therapy based on MRD response may increase the number of AML patients able to therapy with curative intent, currently many of those diagnosed do not
receive best available therapy\textsuperscript{13} and may benefit from the titration of safe, moderately effective therapy based on the objective high sensitivity response criteria of MRD.

How the results of MRD assessments are ultimately integrated into standard cytogenetic and molecular risk stratification systems remains an open question\textsuperscript{129} that ultimately will only be answered by prospective clinical trials; while there is clear evidence that disease biology in the form of pre-treatment risk stratification factors influence risk of having residual disease after initial treatment\textsuperscript{93} it appears that deep negativity in MRD measurements post-treatment is an independent protective factor against relapse and that this factor may trump other considerations in some\textsuperscript{17} but not all\textsuperscript{95} situations.

Many recent excellent reviews, editorials and comment pieces have advocated for the incorporation of MRD monitoring into AML clinical trials\textsuperscript{20, 22–24, 30, 130–134}. It is our hope that by reviewing the most recent clinical trial results in this area we may add our voices to this chorus. While much work remains to standardize the timing, performance, indications and application of MRD monitoring across multiple centers this should not preclude, in the meantime, the more immediate establishment of local AML MRD monitoring capacity. Given our belief that the current standard of care therapy for patients in the USA diagnosed with acute myeloid leukemia, if treated with curative intent, is referral to a specialized center where an appropriate clinical trial can be offered\textsuperscript{135}, we would add now an additional characteristic of such a leukemia center of excellence would be the capacity to routinely provide MRD monitoring for the majority of cases of AML and to universally integrate such monitoring into all future therapeutic clinical trials.

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References


Nat Rev Clin Oncol. Author manuscript; available in PMC 2014 September 15.

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59. Kristensen T, Moller MB, Friis L, Bergmann OJ, Preiss B. NPM1 mutation is a stable marker for minimal residual disease monitoring in acute myeloid leukaemia patients with increased sensitivity compared to WT1 expression. Eur J Haematol. 2011; 87:400–8. [PubMed: 21707751]


62. Shayegi N, et al. The level of residual disease based on mutant NPM1 is an independent prognostic factor for relapse and survival in AML. Blood. 2013


Nat Rev Clin Oncol. Author manuscript; available in PMC 2014 September 15.


Biographies

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Judith E. Karp is Director of the Leukemia Program at the Johns Hopkins Sidney Kimmel Comprehensive Cancer Center and Professor of Oncology and Professor of Medicine at the Johns Hopkins School of Medicine. She received her medical degree from Stanford University School of Medicine, completed internship and residency at Stanford University Hospital and Johns Hopkins Hospital, and completed her Oncology Fellowship at the Johns Hopkins University School of Medicine. From 1990–1995, she served as Special Assistant to Director, National Cancer Institute. Her research interests focused on the experimental therapeutics of acute leukemias, including development of timed sequential therapy, new biologic agents for older adults with acute leukemias, and new approaches to the treatment of refractory acute leukemias including secondary leukemias that evolve from myelodysplasia or from prior cytotoxic chemotherapies.
Figure 1. Detection thresholds of various MRD modalities compared to traditional clinical complete remission
Axis scales approximate and solely for illustrative purposes.
http://www.nature.com/nrclinonc/journal/v10/n8/fig_tab/nrclinonc.2013.100_F1.html and
http://www.nature.com/nrclinonc/journal/v10/n8/fig_tab/nrclinonc.2013.100_F2.html
### Table 1
Evolution of response criteria for complete remission in adult patients with Acute Myeloid Leukemia.

<table>
<thead>
<tr>
<th></th>
<th>1956</th>
<th>2012</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone Marrow</td>
<td>Less than 5% blasts and absence of cells that can be individually identified as leukemic.</td>
<td>Less than 5% blasts including no blasts with Auer rods.</td>
</tr>
<tr>
<td>Extramedullary disease</td>
<td>Subsidence of all evidence of leukemic infiltration</td>
<td>No residual evidence of extramedullary disease</td>
</tr>
<tr>
<td>Platelets</td>
<td>&gt;100,000</td>
<td>&gt;100,000</td>
</tr>
<tr>
<td>Neutrophil count</td>
<td>&gt;200/mm³</td>
<td>&gt;1000/mcL</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>&gt;12 gm/dl for 1 month</td>
<td>Transfusion-free</td>
</tr>
<tr>
<td>Clinical</td>
<td>No symptoms ascribable to leukemia</td>
<td>No criteria.</td>
</tr>
</tbody>
</table>

An additional category of “cytogenetics normal in those with previously abnormal cytogenetics” is listed but not required for a morphological complete response in the complete remission category of 2012 NCCN guidelines and is “commended primarily for use in clinical research studies” in the current international working group (IWG) criteria that those guidelines are based on. There is evidence however that persistence of cytogenetic abnormalities in AML patients in complete remission is associated with a worse prognosis.
Table 2
Comparison of Flow Cytometry versus PCR for detection of MRD in AML

<table>
<thead>
<tr>
<th>Flow Cytometry</th>
<th>Polymerase Chain Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Requires significant human capital as highly trained pathologist expertise required for interpretation.</td>
<td>Can be run in any certified laboratory with real-time PCR capacity including off-site central reference laboratory.</td>
</tr>
<tr>
<td>Results within one day</td>
<td>Results may take several days</td>
</tr>
<tr>
<td>Single result interpretable</td>
<td>Interpretation often requires trend of results</td>
</tr>
<tr>
<td>Excellent specificity when using defined LAIP</td>
<td>Low level transcripts in remission samples common – requires setting of threshold limits.</td>
</tr>
<tr>
<td>Requires use multiple LAIPs, multiple antibody flurochrome combinations and at least 200,000 events for adequate sensitivity.</td>
<td>Excellent sensitivity, at least equal and in some cases superior to flow cytometry.</td>
</tr>
<tr>
<td>Leukemic phenotype may “change” over time, initial LAIP may not identify subclones responsible for relapse.</td>
<td>Many targets stable</td>
</tr>
<tr>
<td>Allows detection of cells with leukemia stem cell phenotype.</td>
<td>Does not allow discrimination of “bulk” from “stem cell” leukemia.</td>
</tr>
<tr>
<td>Detection to $10^{-4}$ possible</td>
<td>Detection from $10^{-4}$ to $10^{-6}$ possible</td>
</tr>
</tbody>
</table>
### Table 3
Recent informative trials and reports regarding use of MRD in AML

<table>
<thead>
<tr>
<th>Trial</th>
<th>Year Reported</th>
<th>AML subtype</th>
<th>MRD Detection Method</th>
<th>Key Points</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>AML02</td>
<td>2010 2012</td>
<td>Childhood AML</td>
<td>FC, Mut-PCR</td>
<td>Multicenter trial. FC-based MRD–directed therapy achieved impressive outcomes in 216 children with AML.</td>
<td>56, 57</td>
</tr>
<tr>
<td>EORTC/GIMEMA</td>
<td>2010</td>
<td>Adult AML</td>
<td>FC</td>
<td>Retrospective analysis of 123 patients. Used MRD status at end of consolidation together with pre-treatment cytogenetic and FLT3 to segregate into low and high relapse risk groups.</td>
<td>95</td>
</tr>
<tr>
<td>AMLSG</td>
<td>2011</td>
<td>Adults, NPM1mut AML</td>
<td>Mut-PCR</td>
<td>NPM1mut seen in 25–30% of cases of AML. This report of 245 patients with NPM1mut showed transcript levels was strongly associated with relapse risk and prognosis after each treatment. Demonstrated PB sufficient for monitoring during follow-up.</td>
<td>58</td>
</tr>
<tr>
<td>MRC AML-15</td>
<td>2012</td>
<td>Adult CBF-AML</td>
<td>Mut-PCR</td>
<td>278 patients. RT-PCR for CBF AML transcripts allowed establishment of thresholds associated with 100% risk of relapse. Demonstrated PB sufficient for monitoring during follow-up.</td>
<td>16</td>
</tr>
<tr>
<td>Kwon et al.</td>
<td>2012</td>
<td>Adult AML</td>
<td>WT1-PCR</td>
<td>In a series of 21 AML patients followed after SCT, neither FC nor chimerism outperformed WT1 RQ-PCR for relapse anticipation.</td>
<td>52</td>
</tr>
<tr>
<td>Rossi et al.</td>
<td>2012</td>
<td>Adult AML</td>
<td>WT1-PCR</td>
<td>In 23 AML patients followed by MRD monitoring, six color FC was comparable to WT1 RQ-PCR technically and in clinical significance.</td>
<td>51</td>
</tr>
<tr>
<td>Yan et al.</td>
<td>2012</td>
<td>AML, AML, MDS Age 2–59</td>
<td>FC, WT1-PCR</td>
<td>Non-ranomized retrospective review of post-SCT experience at Peking University. Development of MRD after SCT associated with increased relapse risk, but less relapse observed in those who received modified DLI therapy.</td>
<td>115</td>
</tr>
<tr>
<td>RELAZA</td>
<td>2012</td>
<td>Adult AML/MDS</td>
<td>CD34+ donor chimerism</td>
<td>Open label single center phase II. Azacitidine intervention for development of MRD after SCT can on average delay, but not always prevent, hematological relapse.</td>
<td>113</td>
</tr>
<tr>
<td>COG AAML03P1</td>
<td>2012</td>
<td>AML (aged under 21)</td>
<td>FC</td>
<td>Presence of FC defined MRD (seen in 25% of patients) while in CR after first induction (EOI1) was highly correlated with relapse. MRD at EOI1 was correlated with cytogenetic/molecular risk.</td>
<td>93</td>
</tr>
<tr>
<td>CBF-2006</td>
<td>2013</td>
<td>Adult CBF-AML</td>
<td>Mut-PCR</td>
<td>198 patients. Multicenter trial. RQ-PCR for CBF transcript MRD, but not KIT or FLT3 mutation status, was prognostic of relapse.</td>
<td>17</td>
</tr>
<tr>
<td>AML05</td>
<td>2013</td>
<td>t(8,21) AML Aged 15–60</td>
<td>Mut-PCR</td>
<td>MRD after consolidation cycle 2 predictive of relapse risk; those at this high risk had better RFS and OS if received allo-SCT.</td>
<td>116</td>
</tr>
</tbody>
</table>
### Table 4

Clinical scenarios in the care of AML patients where MRD may have utility.

- High-resolution determination of efficacy of therapy
- To allow target-driven titration of dose and duration of treatment
- Relapse risk stratification after induction to allow triage to optimal consolidation therapy
- To determine prognosis after completion of standard treatment
- To spare toxicity and cost of SCT in those with low risk of relapse
- Assignment to maintenance therapy after completion of standard treatment