Open Access Full Text Article

ORIGINAL RESEARCH

Use of FLT3 inhibitors in acute myeloid leukemia remission induction or salvage therapy: systematic review and meta-analysis

Minglei Yang* Jian Zhao* Tielong Liu Xinghai Yang Haifeng Wei Wei Xu Jianru Xiao

Department of Orthopedic Oncology, Changzheng Hospital, Second Military Medical University, Shanghai, 200003, China

*These authors contributed equally to this work

Correspondence: Wei Xu; Jianru Xiao Department of Orthopedic Oncology, Changzheng Hospital, Second Military Medical University, 415 Fengyang Road, Shanghai 200003, China Tel +86 21 137 6127 8657; +86 21 137 0178 5283 Fax +86 21 8188 5634; +86 21 8188 5634 Email xuweichangzheng@hotmail.com; jianruxiao83@163.com



Background: Previous studies showed that FLT3 inhibitors played an important role in acute myeloid leukemia (AML) therapy. However, discrepancies remain regarding the association between FLT3 inhibitors use and prognosis of AML patients in clinical trials.

Aim: The aim of this study was to evaluate the effect of FLT3 inhibitors on the treatment of AML in a systematic review and meta-analysis.

Materials and methods: PubMed, Embase, and Cochrane Library databases were searched for studies published before August 2017 that used FLT3 inhibitors in AML. Fixed- and random-effect models were used, and between-study heterogeneity was assessed.

Results: A total of 26 studies fitting our inclusion and exclusion criteria were included. The FLT3 status of patients and main treatment outcomes including overall survival (OS), event-free survival (EFS), relapse-free survival (RFS), complete remission (CR), and overall response rate (ORR) after therapy were extracted. Five studies comparing addition of FLT3 inhibitors and placebo or blank control to chemotherapy were analyzed in Part I, showing improved OS (hazard ratio [HR]=0.86, 95% confidence interval [CI]=0.75–0.99, P=0.03) in the FLT3 inhibitor group but without a significant improvement on EFS (HR=0.86, 95% CI=0.62–1.21, P=0.39) and ORR (odds ratio [OR]=1.10, 95% CI=0.89–1.35, P=0.38). Twenty-one studies evaluating the benefit of using FLT3 inhibitors in different FLT3-type AML patients were more sensitive to FLT3 inhibitor treatment and achieved better CR (OR=1.89, 95% CI=1.06–3.37, P=0.03) and ORR (OR=3.07, 95% CI=2.13–4.43, P<0.001).

Conclusion: Our study showed that combined use of FLT3 inhibitors improved OS and that the FLT3 status of AML patients could affect their sensitivity to FLT3 inhibitors in terms of CR and ORR.

Keywords: acute myeloid leukemia, FLT3-ITD, clinical outcome, meta-analysis

Introduction

Acute myeloid leukemia (AML) is a heterogeneous hematologic malignancy characterized by clonal expansion of myeloid blasts in the peripheral blood, bone marrow (BM), and/or other tissues, with an incidence of 3.7/100,000 individuals.^{1,2} Patients with AML are divided into different risk states based on their cytogenetic and molecular abnormalities. FLT3 protein is a member of the receptor-tyrosine kinase (RTK) family, playing an important role in normal growth and differentiation of hematopoietic precursor cells.^{3,4} In currently available studies,^{4,5} the internal tandem duplications (ITDs) in FLT3 mutations were related to the blast of leukemia cells

Construction of the full terms of this license are available at https://www.dovepress.com/terms. by and incorporate the (retarive Commons Attribution — Non Commercial (unported, v3.0) License (http://creative.commons.org/license/by-nc/3.0). By accessing the work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs 4.2 and 5 of our Terms (https://www.dovepress.com/terms.php). and suppression of apoptosis and were detected in ~23% AML patients. A meta-analysis⁶ of AML patients showed that the pooled hazard ratio (HR) of the impact of FLT3–ITD on overall survival (OS) and relapse-free survival (RFS) was 1.86 (95% confidence interval [CI]=1.57–2.20) and 1.83 (95% CI=1.54–2.18), respectively. The goal of treatment of AML is to achieve morphologic, cytogenetic, and molecular complete remission (CR); prepare for subsequent consolidation chemotherapy or hematopoietic cell transplantation (HCT); and get long-term OS, event-free survival (EFS), and RFS.² Patients with FLT3–ITD mutations are considered to a have a higher risk of relapse with shorter

RFS, EFS, and OS and are recommended for clinical trials

where available.5,7 FLT3 inhibitors are a type of tyrosine kinase inhibitors with FLT3 inhibitory activity. Preclinical research and clinical trials^{8,9} have confirmed the activity of FLT3 inhibitors against AML primary blasts with FLT3-ITD mutations in vitro and in vivo. Current studies^{8,10} have demonstrated that the patients can benefit from the use of FLT3 inhibitors. Sunitinib, midostaurin, lestaurtinib, sorafenib, quizartinib, and crenolanib are the most common FLT3 inhibitors reported in current clinical trials. Most of these drugs were first approved for use in solid malignancies by the US Food and Drug Administration (FDA). Except for several Phase I and Phase II clinical trials on the safety and effectiveness of these compounds,¹¹⁻³¹ there have been limited studies to evaluate the dosage, duration, monotherapy, or combination use with chemotherapy of these drugs in AML.³² In addition, the demographic baselines in these studies varied substantially. Only a few randomized controlled trials (RCTs) or Phase III clinical trials^{33–37} have compared midostaurin-, lestaurtinib-, or sorafenib-containing new regimens with the standard chemotherapy regimens.

There is no consensus on the role of FLT3 inhibitors in AML patients. The outcomes of clinical trials are controversial, and high risks of adverse events are also associated with these agents.³³ Knowing that there is no systematic review and meta-analysis focusing on the role of FLT3 inhibitors in AML therapy, we conducted this meta-analysis in an attempt to clarify the clinical benefit of FLT3 inhibitors and make a summary on the clinical outcomes in terms of CR, overall response rate (ORR), OS, and EFS when FLT3 inhibitors are added to conventional chemotherapy by comparing them with chemotherapy alone (Part I). We also compared CR or ORR between AML patients with different FLT3–ITD statuses when FLT3 inhibitors were used (Part II).

Materials and methods Search strategy

The following literature searching strategy was used to identify potentially relevant studies focusing on FLT3 inhibitors in AML induction/salvage therapy. PubMed, Embase, and Cochrane Library databases were searched by using the following index terms: leukemia, myeloid, acute AND FLT3 OR tyrosine kinase inhibitor OR sorafenib OR sunitinib OR midostaurin OR lestaurtinib OR quizartinib OR crenolanib, without restrictions on the region and publication type. Mesh subject words with random words were selected from the PubMed database. No language limit was required for publication. Relevant reference lists were screened through manual search for further potential studies. When multiple publications reported on the same study population, the report that was most complete or had the longest follow-up period was used. The last date of the search was August 3, 2017 (Figure 1).

Study selection

Two reviewers independently reviewed all studies and selected eligible trials. To analyze the potential benefit from adding FLT3 inhibitors to AML treatment in mixed population, we used the following inclusion criteria in Part I: 1) RCTs involving comparison between FLT3 inhibitors and placebo or blank control, with or without other chemotherapy compounds in AML remission induction and/or salvage treatment; 2) studies using CR, morphologic CR with incomplete blood count recovery (CRi), morphologic CR with incomplete platelet recovery (CRp), partial remission (PR), OS, RFS, and EFS as the response criteria; and 3) the definition of the above response criteria in accordance with the recommendations of the International Working Group.³⁸ CR was defined by the presence of <5% blasts in the BM with 1×10^{9} /L neutrophils and 100×10^{9} /L platelets in the peripheral blood. CRp was defined in patients with CR but persistent platelet count <100×109/L. CRi was defined in patients with persistent neutrophil count $<1\times10^{9}/L$ or without platelet recovery. PR required all hematologic values for a CR but with a decrease of at least 50% in the percentage of blasts to 5-25% in the BM aspirate. To analyze the possible effect of FLT3 inhibitors on AML with different FLT3 statuses, we used the following inclusion criteria in Part II: 1) clinical trials involving the outcome of using FLT3 inhibitors for the treatment of AML; 2) studies using CR, CRi, CRp, PR, OS, RFS, and EFS as the response criteria; 3) patients involving gene analysis and treatment outcomes stratified on different



Figure I Flow diagram concerning study identification, inclusion, and exclusion.

Abbreviations: AML, acute myeloid leukemia; FLT3, FMS-like tyrosine kinase 3; WT, wild type; NO, without.

FLT3 status; and 4) studies involving patients with multiple types of FLT3 status. Owing to high AML transformation in myelodysplastic syndrome (MDS) and related therapeutic regimens between AML and MDS, studies with fewer than 10% MDS patients were included in our analysis. Studies that could not meet any one of the above inclusion criteria were excluded. Animal studies, review articles, and case reports were also excluded.

Data extraction

Two reviewers independently extracted the data from the full manuscripts of included articles, and any disagreement was resolved by the third reviewer. We obtained the following information from each article: first author, year of publication, number of included patients, gene type of patients, therapeutic regimen, prior therapeutic regimen, type of AML, AML status, and treatment outcome. The main treatment outcomes included CR, CRi, CRp, PR, OS, RFS, and EFS after AML induction or salvage therapy in different regimens (in Part I) or in different FLT3 status patients (in Part II). Patients with the therapy outcome better than or equal to PR were thought to respond to the current treatment regimen and calculated for ORR. The quality of studies was assessed by using the Cochrane risk of bias assessment tool for RCTs (Figure S1) and Newcastle–Ottawa Quality Assessment Scale (NOS) for cohort studies (Tables 1 and S1).

Statistical analysis

This meta-analysis was performed with the aid of software Review Manager Version 5.3 (available at <u>http://ims.cochrane.</u> <u>org/revman/download</u>). The number of main outcome events and simple size were calculated. In the forest plots, odds ratio (OR), HR, and their 95% CI were displayed in our qualitative data synthesis. To assess the heterogeneity between studies,

status status status status status mediar (angle status) Alvarado Y ¹ Na 2014 Multiple types NA Monotherapy Prior treatments 54 (18-87) Coorses [P ¹ NCT01411267 2016 Quizarinib Phase I Chemotherapy Relapsed Ant 521 Correst [P ¹ NCT00462761 2013 Quizarinib Phase I Monotherapy Relapsed refractory Ant 603 Correst [P ¹ NA 2005 Sonitoibi Phase I Monotherapy Relapsed refractory Ant 603 Fedler W ¹ NA 2005 Sonitoibi Phase I Monotherapy Relapsed refractory Ant 71 (37-82) Fedler W ¹ NA 2005 Sonitoibi Phase I Monotherapy Relapsed refractory Ant 71 (37-82) Fedler W ¹ NA 2001 Sonfenib Phase I Monotherapy Relapsed refractory Ant 71 (37-82) Kaapper S ² NA 2010 Sonfenib Phase I Sonfenib NA 2054-80	Author	Clinical register	Year	Suberoup	Clinical	Treatment	Patient states	Age (vears).	Size
Avarado Y1 NA Zoll Avarado Y1 NCT0(111267 Zoll Guizartinib Phase I Monotherapy Relapsed/refractory AVI Zoll Solutinib Phase I Monotherapy quizartinib Relapsed/refractory AVI Zoll Solutinib Phase I Monotherapy quizartinib Relapsed/refractory AVI Zoll Solutinib Phase I Monotherapy Relapsed/refractory AVI Solutinib Phase I Solutinib Phase II Constantio Phase II Solutinib Phase III Solutinib		D		-	status			median (range)	
Cooper TM ¹¹ NCT01411267 2016 Quizartinib Phase I Chemotherapy quizartinib Relapsed/refractory AML 261 Corres JE ¹⁰ NA 2013 Quizartinib Phase I Monotherapy Relapsed/refractory AML 560 Corres JE ¹⁰ NA 2013 Quizartinib Phase I Monotherapy Relapsed/refractory AML 560 Curres JE ¹⁰ NA 2005 Suntinib Phase I Monotherapy Relapsed/refractory AML 60 (32-86) Fielder W ¹⁰ NA 2005 Suntinib Phase II Monotherapy Relapsed/refractory AML 71 (37-82) Fielder W ¹⁰ NA 2005 Suntinib Phase II Monotherapy Relapsed/refractory AML 72 (54-80) Fielder W ¹⁰ NA 2011 Sorafenib Phase III Chemotherapy Relapsed/refractory AML 73 (57-82) Fielder W ¹⁰ NA 2001 Sorafenib Phase III Chemotherapy Relapsed/refractory AML 231 (57-82) Maconald DA ¹¹ NA 2012 <td< td=""><td>Alvarado Y¹²</td><td>NA</td><td>2014</td><td>Multiple types</td><td>AA</td><td>Monotherapy</td><td>Prior treatments</td><td>54 (18–87)</td><td>69</td></td<>	Alvarado Y ¹²	NA	2014	Multiple types	AA	Monotherapy	Prior treatments	54 (18–87)	69
Cortes JP ¹⁰ NA 2012 Quizartinib Phase II Monotherapy Relapsed/refractory AML 260 Curmp M ¹⁵ NCT00462761 2013 Quizartinib Phase I Monotherapy Relapsed/refractory AML 60 (33-66) Curmp M ¹⁵ NA 2010 Soratemib Phase I Monotherapy Relapsed/refractory AML 60 (33-66) Fielder W ¹⁴ NA 2005 Sunitinib Phase II Monotherapy Relapsed/refractory AML 60 (33-66) Fielder W ¹⁴ NCT00783553 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML 73 (57-82) Fielder W ¹⁴ NA 2001 Sorafenib Phase II Monotherapy Relapsed/refractory AML 73 (57-82) Knapper S ²⁰ NA 2013 Sorafenib Phase II Monotherapy Relapsed/refractory AML 73 (57-82) Knapper S ²⁰ NA 2013 Sorafenib Phase II Monotherapy Relapsed/refractory AML 73 (57-82) Knapper S ²⁰ NA Z013	Cooper TM ¹³	NCT01411267	2016	Quizartinib	Phase I	Chemotherapy, quizartinib	Relapsed AML	≤21	14
Cortes JE ⁴ NCT00462761 2013 Quizartinib Phase I Monotherapy Relapsed/refractory AML 60 (23-86) Fiedler W ¹ NA 2010 Sorafenib Phase I Monotherapy Relapsed/refractory AML 60 (23-86) Fiedler W ¹ NA 2010 Sorafenib Phase II Monotherapy Relapsed/refractory AML 71 (37-82) Fiedler W ¹ NCT0038533 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML 73 (57-80) Fischer T ¹ s NCT00345942 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML 73 (57-80) Fischer T ¹ s NCT00345942 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML 73 (57-80) Kmapper S ²⁰ NA 2012 Quizartinib Phase II Monotherapy Relapsed/refractory AML 73 (57-80) Kmapper S ²⁰ NA 2013 Sorafenib Phase II Monotherapy Relapsed/refractory AML 73 (57-80) Kmapper S ²⁰ NA	Cortes JE ³⁰	NA	2012	Quizartinib	Phase II	Monotherapy	Relapsed/refractory AML	≥60	133
Crump M ³ NA 2010 Sorafenib Phase I Monotherapy Relapsed/refractory AML or MDS 71 (37-82) Fiedler W ¹ NA 2010 Sunitinib Phase I Monotherapy Relapsed/refractory AML 750 Fiedler W ¹ NA 2015 Sunitinib Phase I Monotherapy Relapsed/refractory AML 260 Fischer T ^{1a} NCT00045942 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML 261 Fischer T ^{1a} NCT00045942 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML 261 Rinaba H ¹⁹ NA 2012 Quartinib Phase II Monotherapy Relapsed/refractory AML 71 (37-82) Kinsple 2013 Sorafenib Phase II Monotherapy Relapsed/refractory AML 73 (67-82) Macdonald DA ²¹ ND.186 2013 Sorafenib Phase I/I Sorafenib Phase I/I Sorafenib Phase I/I Sorafenib Phase I/I Sorafenib PhaseI/I Sorafen	Cortes JE ¹⁴	NCT00462761	2013	Quizartinib	Phase I	Monotherapy	Relapsed/refractory AML	60 (23–86)	54
Fiedler W ¹ NA 2005 Sunitinib Phase I Monotherapy Relapsed/refractory AML 72 (54-80) Fiedler W ¹⁶ NCT00783553 2015 Sunitinib Phase I/I Chemotherapy sunitinib Untreated newly diagnosed AML 260 Fischer T ^{14,a} NCT00783553 2010 Monotherapy Rehapsed/refractory AML 260 Fischer T ^{14,a} NCT00045942 2010 Monotherapy Rehapsed/refractory AML 261 Inaba H ¹⁹ NA 2012 Curatinib Phase I Monotherapy Rehapsed/refractory AML 71(-82) Knapper S ²⁰ NA 2012 Quizartinib Phase I Monotherapy 261 271 Macdonald DA ²¹ ND.186 2013 Sorafenib N Monotherapy 260 231 60 231 60 231 60 60 60 60 65 63 63 66 65 63 66 65 63 63 66 65 63 60 65 63 <t< td=""><td>Crump M¹⁵</td><td>NA</td><td>2010</td><td>Sorafenib</td><td>Phase I</td><td>Monotherapy</td><td>Relapsed/refractory AML or MDS</td><td>71 (37–82)</td><td>27</td></t<>	Crump M¹⁵	NA	2010	Sorafenib	Phase I	Monotherapy	Relapsed/refractory AML or MDS	71 (37–82)	27
Fiedler W ¹⁶ NCT00783653 2015 Sunitinib Phase I/I Chemotherapy, sunitinib Untreated newly diagnosed AML >60 Fischer T ^{18,1} NCT00045942 2010 Midostaurin Phase II Monotherapy, sunitinib Untreated newly diagnosed AML >60 Fischer T ^{18,1} NCT00045942 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML or MDS >18 Knapper S ³⁰ NA 2006 Lestaurtinib Phase II Monotherapy Relapsed/refractory AML 73 (67–82) Levis MJ ¹³ NA 2012 Quizartinib Phase II Monotherapy Relapsed/refractory AML FtT3-ITD (-): Macdonald DA ³¹ ND-I86 2013 Sorafenib N Monotherapy Relapsed/refractory AML FtT3-ITD (-): Macdonald DA ³¹ ND-I86 Phase I/I Sorafenib, and vortanbine No prior chemotherapy >60 Parz KW ¹¹ NA NO Monotherapy Relapsed/refractory AML 53 (8-66) Macdonald DA ³¹ ND-I86 Phase I/I Sorafenib, dataurin, dataubicin, and cytarabine No prior chemotherapy 260 R	Fiedler W ¹⁷	NA	2005	Sunitinib	Phase I	Monotherapy	Relapsed/refractory AML	72 (54–80)	12
Fischer T ^{4a} NCT00045942 2010 Midostaurin Phase II Monotherapy Relapsed/refractory AML or MDS ≥18 Inaba H ^{1a} NA 2011 Sorafenib Phase I Sorafenib, dofarabine, and cytarabine Relapsed/refractory AML or MDS ≥18 Knapper S ²⁰ NA 2006 Lestaurtinib Phase II Monotherapy Untreated AML 73 (67–82) Levis MJ ²⁹ NA 2012 Quizartinib Phase II Monotherapy Untreated AML 73 (67–82) Levis MJ ²⁹ NA 2012 Quizartinib Phase II Monotherapy Nonotherapy Relapsed/refractory AML 73 (67–82) Macdonald DA ²¹ IND.186 2013 Sorafenib N Monotherapy Relapsed/refractory AML 73 (67–82) Macdonald DA ²¹ IND.186 2013 Sorafenib N Monotherapy and midostaurin Relapsed/refractory AML 73 (67–82) Macdonald DA ²¹ ND.186 N Monotherapy and midostaurin No prior chemotherapy McIan 63 Rausingh G ²² NCT01161550 2014 Midostaurin, themotherapy and midostaurin Na Pased/refractory AML	Fiedler W ¹⁶	NCT00783653	2015	Sunitinib	Phase I/II	Chemotherapy, sunitinib	Untreated newly diagnosed AML	≥60	22
Inda H1 ⁹ NA 2011 Sorafenib Phase I Sorafenib, clofarabine, and cytarabine Relapsed/ refractory AML ≤21 Knapper S ²⁰ NA 2006 Lestaurtinib Phase II Monotherapy Untreated AML 73 (57–82) Levis MJ ²⁹ NA 2012 Quizartinib Phase II Monotherapy Untreated AML 73 (57–82) Levis MJ ²⁹ NA 2013 Sorafenib Phase II Monotherapy Relapsed/refractory AML 73 (57–82) Macdonald DA ²¹ IND.186 2013 Sorafenib N Monotherapy Relapsed/refractory AML 73 (57–82) Prazt KW ¹¹ NA 2010 Sorafenib N Monotherapy and midostaurin Relapsed/refractory AML 52 (32–71) Ravandi F ^{24b} NA 2014 Midostaurin Phase I/I Sorafenib, cytarabine Untreated AML 53 (18–66) Ravandi F ^{24b} NA 2014 Sorafenib Phase I/I Sorafenib, cytarabine Untreated AML 53 (18–66) Ravandi F ^{24b} NA 2013 Sorafenib Phase I/I Sorafenib, cytarabine, und cytarabine Untreated AM	Fischer T ^{18,a}	NCT00045942	2010	Midostaurin	Phase IIB	Monotherapy	Relapsed/refractory AML or MDS	≥I8	92
Knapper S ²⁰ NA 2006 Lestaurtinib Phase II Monotherapy Untreated AML 73 (67–82) Levis MJ ²⁹ NA 2012 Quizartinib Phase II Monotherapy Relapsed/refractory AML FLT3-ITD (-): Macdonald DA ²¹ IND.186 2013 Sorafenib Phase I/I Sorafenib and low-dose cytarabine No prior chemotherapy 260 Macdonald DA ²¹ IND.186 2013 Sorafenib N Monotherapy Relapsed/refractory AML FLT3-ITD (-): Macdonald DA ²¹ IND.186 2010 Sorafenib N Monotherapy Relapsed/refractory AML 76.7-30 Ramsingh G ²² NCT01161550 2014 Midostaurin Phase I/I Sorafenib, idarubicin, and cytarabine No prior chemotherapy 260 Ravandi F ^{24,b} NA 2016 Sorafenib, cytarabine, and idarubicin, and cytarabine Untreated AML 52 (32–71) Ravandi F ^{24,b} NA 2014 Sorafenib, cytarabine, and idarubicin, and cytarabine Untreated AML 53 (18–66) Stone RM ^{36,c} NA 2012 Sorafenib Phase II Sorafenib, cytararabine, and idarubicin Untrea	Inaba H ¹⁹	NA	2011	Sorafenib	Phase I	Sorafenib, clofarabine, and cytarabine	Relapsed/ refractory AML	≤21	=
Levis MJ ²³ NA 2012 Quizartinib Phase II Monotherapy Relapsed/refractory AML FLT3-ITD (+): Macdonald DA ¹¹ ND:186 2013 Sorafenib Phase I/II Sorafenib and low-dose cytarabine No prior chemotherapy >60 Prazz KW ¹¹ NA 2010 Sorafenib N Monotherapy additional 260 Prazz KW ¹¹ NA 2010 Sorafenib N Monotherapy additional 260 Ransingh G ²² NCT01161550 2014 Midostaurin Phase I/II Sorafenib, idarubicin, and cytarabine No prior chemotherapy 260 Ravandi F ²⁴ b NA 2010 Sorafenib Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML 55 (32-71) Ravandi F ²⁴ b NA 2012 Midostaurin Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML 55 (32-71) Stone RM ³⁶⁴ NA 2012 Midostaurin, Phase II Sorafenib, idarubicin, and cytarabine Untreated AML 56 (32-90) Stone RM ³⁶⁴ NA 2012 Midostaurin, Phase II Sorafenib, idarubicin, and cytarabine <td< td=""><td>Knapper S²⁰</td><td>NA</td><td>2006</td><td>Lestaurtinib</td><td>Phase II</td><td>Monotherapy</td><td>Untreated AML</td><td>73 (67–82)</td><td>27</td></td<>	Knapper S ²⁰	NA	2006	Lestaurtinib	Phase II	Monotherapy	Untreated AML	73 (67–82)	27
Macdonald DA ²¹ IND.186 2013 Sorafenib Phase I/I Sorafenib No Phase I/I Sorafenib No Pisse I/I Sorafenib No Monotherapy Relapsed/refractory AML Pielon 63 Ravandi F ²¹ NA 2010 Sorafenib N Monotherapy and midostaurin Relapsed/refractory AML Median 63 Ravandi F ²¹ NA 2010 Sorafenib Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML 52 (32-71) Ravandi F ²¹ NA 2014 Sorafenib Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML 53 (18-66) Stone RM ^{3dc} NA 2012 Midostaurin Phase I Midostaurin, bortezomib, and MEC Relapsed/refractory AML 53 (18-66) Stone RM ^{3dc} NA 2012 Midostaurin Phase I Midostaurin, bortezomib, and MEC Relapsed/refractory AML 53 (18-66) Walker AR ^{3dd} NCT0117488 2016 Midostaurin, bortezomib, and ME	Levis MJ ²⁹	NA	2012	Quizartinib	Phase II	Monotherapy	Relapsed/refractory AML	FLT3-ITD (+): 50 (19-77)	137
Macdonald DA ²¹ IND.186 2013 Sorafenib Phase I/I Sorafenib and low-dose cytarabine No prior chemotherapy ≥60 Prazt KW ¹¹ NA 2010 Sorafenib N Monotherapy Relapsed/refractory AML ≥60 Ramsingh G ²² NCT01161550 2014 Midostaurin Phase I Chemotherapy and midostaurin Relapsed/refractory AML 52 (32-71) Ravaidi F ²⁴ NA 2010 Sorafenib Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML 53 (32-71) Ravaidi F ²⁴ NA 2010 Sorafenib Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML 53 (32-71) Ravaidi F ²³ NA 2012 Sorafenib Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML 53 (32-71) Ravaidi F ²³ NA 2012 Midostaurin Phase I/I Sorafenib, idarubicin Untreated AML 53 (18-66) Stone RM ^{36c} NA 2012 Midostaurin, Phase I Midostaurin, bortezomib, and MEC Relapsed/refractory AML 19-70 Valker AR ^{38d} NCT0117488 2016 Midostaurin, bortezomib, and M								FLT3-ITD (-): 55 (30-73)	
Prazz KW ^{II} NA 2010 Sorafenib N Monotherapy Relapsed/refractory AML Median 63 Ramsingh G ²² NCT01161550 2014 Midostaurin Phase I Chemotherapy and midostaurin Relapsed/refractory AML 52 (32–71) Ravandi F ^{24.b} NA 2010 Sorafenib Phase I Chemotherapy and midostaurin Relapsed/refractory AML 53 (32–71) Ravandi F ^{24.b} NA 2010 Sorafenib Phase II Sorafenib, idarubicin and cytarabine Untreated AML 55 (32–71) Ravandi F ^{24.b} NA 2014 Sorafenib Phase II Sorafenib, idarubicin Untreated AML 53 (32–60) Stone RM ³⁶⁴ NCT0117488 2016 Midostaurin, Phase IB Midostaurin, chemotherapy Untreated AML 53 (18–66) Kalker AR ³⁶⁴ NCT01174888 2016 Midostaurin, bortezomib, and MEC Relapsed/refractory AML 19–70 Kalker AR ³⁶⁴ NCT01174888 2016 Midostaurin, bortezomib, and MEC Relapsed/refractory AML 19–70 Kalker AR ³⁶⁴ NCT01174888 2016 Crenolanib NA Mixed 65.5 (19–82)	Macdonald DA ²¹	IND.186	2013	Sorafenib	Phase I/II	Sorafenib and low-dose cytarabine	No prior chemotherapy	≥60	21
Ramsingh G22NCT011615502014MidostaurinPhase IChemotherapy and midostaurinRelapsed/refractory AML52 (32–71)Ravandi F24NA2010SorafenibPhase I/ISorafenib, idarubicin, and cytarabineUntreated AML55Ravandi F23NA2014SorafenibPhase IISorafenib, idarubicinUntreated AML565Stone RM ^{24c} NA2012MidostaurinPhase IISorafenib, cytarabine, and idarubicinUntreated AML53 (18–66)Stone RM ^{24c} NA2012MidostaurinPhase IBMidostaurin, chemotherapyUntreated AML53 (18–66)Stone RM ^{24c} NA2012MidostaurinPhase IBMidostaurin, bortezomib, and MECRelapsed/refractory AML19–70Solig C ³³ NA2012SorafenibNAMixed64.5 (19–82)Valker AR ^{34d} NA2012SorafenibNAMixed64.5 (19–82)Ver SP ³¹ NA2016CrenolanibNAMixed64.5 (19–82)Ver SP ³¹ NA2016CrenolanibNA201670.5 (22–90)Mater AR ^{34d} A2016CrenolanibNA201620-00Ver SP ³¹ NA2016CrenolanibNA201620-00Ver SP ³¹ NA2016CrenolanibNA201620-00Ver SP ³¹ NA2016CrenolanibNA201620-00Ver SP ³¹ NA2016CrenolanibNA20-00 <t< td=""><td>Pratz KW^{II}</td><td>NA</td><td>2010</td><td>Sorafenib</td><td>z</td><td>Monotherapy</td><td>Relapsed/refractory AML</td><td>Median 63</td><td>12</td></t<>	Pratz KW ^{II}	NA	2010	Sorafenib	z	Monotherapy	Relapsed/refractory AML	Median 63	12
Ravandi F ^{24b} NA 2010 Sorafenib Phase I/I Sorafenib, idarubicin, and cytarabine Untreated AML ≤65 Ravandi F ²³ NA 2014 Sorafenib Phase II Sorafenib, cytarabine, and idarubicin Untreated AML 53 (18–66) Stone RM ^{2ac} NA 2012 Midostaurin Phase II Sorafenib, cytarabine, and idarubicin Untreated AML 53 (18–66) Stone RM ^{2ac} NA 2012 Midostaurin Phase IB Midostaurin, chemotherapy Untreated AML 53 (18–66) Walker AR ^{3ad} NCT01174888 2016 Midostaurin, phase IB Midostaurin, bortezomib, and MEC Relapsed/refractory AML 19–70 Röllig C ³⁵ NA 2012 Sorafenib NA Mixed 64.5 (19–82) Ver Sp ¹¹ NA 2016 Crenolanib NA Mixed 64.5 (19–82) Ver Sp ¹¹ NA 2016 Crenolanib NA Mixed 64.5 (19–82) Ver Sp ¹¹ NA 2016 Crenolanib, salvage chemotherapy, Prior treatments 70.5 (22–90) Mathematical AML AM AM AM AM	Ramsingh G ²²	NCT01161550	2014	Midostaurin	Phase I	Chemotherapy and midostaurin	Relapsed/refractory AML	52 (32–71)	6
Ravandi F2NA2014SorafenibPhase IISorafenib, cytarabine, and idarubicinUntreated AML53 (18–66)Stone RM ^{3Ac} NA2012MidostaurinPhase IBMidostaurin, chemotherapyUntreated AMLFigure 48.5Stone RM ^{3Ac} NA2016MidostaurinPhase IBMidostaurin, bortezomib, and MECRelapsed/refractory AML19–70Walker AR ^{3ad} NCT011748882016MidostaurinPhase IMidostaurin, bortezomib, and MECRelapsed/refractory AML19–70Köllig C ³⁵ NA2012SorafenibNAMixed64.5 (19–82)Ver Sp ³¹ NA2016CrenolanibNACrenolanib, salvage chemotherapy,Prior treatments70.5 (22–90)MaxMax2016CrenolanibNAMixed64.5 (19–82)Ver Sp ³¹ NA2016CrenolanibNA70.5 (22–90)MaxMaxMAMAMA70.5 (22–90)MaxMaxMAMAMA70.5 (22–90)MaxMaxMAMAMA70.5 (22–90)MaxMaxMAMAMA70.5 (22–90)MaxMaxMAMAMA70.5 (22–90)MaxMaxMAMA70.5 (22–90)MaxMaxMAMA70.5 (22–90)MaxMaxMAMA70.5 (22–90)MaxMaxMAMA70.5 (22–90)MaxMaxMA70.5 (22–90)MaxMax	Ravandi F ^{24,b}	NA	2010	Sorafenib	Phase I/II	Sorafenib, idarubicin, and cytarabine	Untreated AML	≤65	51
Stone RM ^{24c} NA 2012 Midostaurin Phase IB Midostaurin, chemotherapy Untreated AML Median 48.5 Walker AR ^{28d} NCT01174888 2016 Midostaurin Phase I Midostaurin, bortezomib, and MEC Relapsed/refractory AML 19–70 Röllig C ²⁵ NA 2012 Sorafenib NA Mixed Mixed 64.5 (19–82) Iyer Sp ¹¹ NA 2016 Crenolanib NA Crenolanib, salvage chemotherapy, Prior treatments 70.5 (22–90) and HAM	Ravandi F ²³	NA	2014	Sorafenib	Phase II	Sorafenib, cytarabine, and idarubicin	Untreated AML	53 (18–66)	62
Walker AR ²⁶⁴ NCT01174888 2016 Midostaurin Phase I Midostaurin, bortezomib, and MEC Relapsed/refractory AML 19–70 Röllig C ²⁵ NA 2012 Sorafenib NA Mixed 64.5 (19–82) Iyer SP ¹¹ NA 2016 Crenolanib NA Crenolanib, salvage chemotherapy, Prior treatments 70.5 (22–90) In And And And And And 70.5 (22–90)	Stone RM ^{26,c}	NA	2012	Midostaurin	Phase IB	Midostaurin, chemotherapy	Untreated AML	Median 48.5	40
Röllig C ²⁵ NA 2012 Sorafenib NA Mixed 64.5 (19-82) Iyer SP ³¹ NA 2016 Crenolanib NA Crenolanib, salvage chemotherapy, Prior treatments 70.5 (22-90) Iver SP ³¹ NA 2016 Crenolanib NA Crenolanib, salvage chemotherapy, Prior treatments 70.5 (22-90)	Walker AR ^{28,d}	NCT01174888	2016	Midostaurin	Phase I	Midostaurin, bortezomib, and MEC	Relapsed/refractory AML	19–70	23
lyer SP ³¹ NA 2016 Crenolanib NA Crenolanib, salvage chemotherapy, Prior treatments 70.5 (22–90) and HAM	Röllig C ²⁵	NA	2012	Sorafenib	AA	Mixed	Mixed	64.5 (19–82)	28
	lyer SP ³¹	NA	2016	Crenolanib	AA	Crenolanib, salvage chemotherapy,	Prior treatments	70.5 (22–90)	6
						and HAM			
Zhang W ²² C I EP25X506801 ZUU8 Soratenib NA Monotherapy Prior treatments 61.3 (48–81)	Zhang W^{27}	CTEP25XS06801	2008	Sorafenib	NA	Monotherapy	Prior treatments	61.5 (48–81)	16

Table I Baseline characteristics of 21 clinical trials included in the meta-analysis (Part 2)

Yang et al

the Cochran's Q test and I^2 test were used. If the Cochran's Q test showed P < 0.05 or the I^2 test exhibited P > 50%, indicating significant heterogeneity between studies, the random-effect model would be used. If the heterogeneity was acceptable, the fixed-effect model would be used. Subgroup analyses on different types of FLT3 inhibitors were performed to further evaluate the heterogeneity and sensitivity. Funnel plots were used to analyze the publication bias.

Results

Study characteristics

A total of 26 studies were identified and included in our study. Five RCTs providing the data about addition of FLT3 inhibitor versus placebo or blank control to conventional chemotherapy in AML were analyzed in Part I. The other 21 articles were analyzed in Part II to evaluate the benefit of using FLT3 inhibitors between AML patients with and without FLT3-ITD mutations. There were 1905 patients involved in Part I and 866 patients involved in Part II. Features of the included studies are summarized in Tables 1 and 2. In Part I, two studies^{33,36} used sorafenib in 464 patients, two^{34,35} used lestaurtinib in 724 patients, and one³⁹ used midostaurin in 717 patients. Four studies^{33,34,36,37} selected untreated AML, and the other³⁵ involved patients with relapsed AML. In Part II, eight studies^{11,15,19,21,23–25,27} used sorafenib, four^{13,14,29,30} used quizartinib, four^{18,22,26,28} used midostaurin, two^{16,17} used sunitinib, one²⁰ used lestaurtinib, one³¹ used crenolanib, and one¹² used multiple types of FLT3 inhibitors. In one study,¹⁸ ORR included not only CR, CRi, CRp, and PR but also hematologic improvement (HI) and blast response (BR).

Part I: effects of adding FLT3 inhibitors to conventional chemotherapy

In five studies included in this part, the end points including OS, EFS, and ORR (containing CR, CRp, and CRi) were analyzed for assessing the clinical effect. HR of OS from four articles (two sorafenib, one lestaurtinib, and one midostaurin) was pooled with no significant heterogeneity between them (P=0.56, I²=0%). Fixed-effect models were selected, and a forest plot was performed to show the results (HR=0.86, 95% CI=0.75-0.99, P=0.03; Figure 2). HR of EFS from three articles (two sorafenib and one midostaurin) was pooled using random-effect models (HR=0.86, 95% CI=0.62-1.21, P=0.39), with heterogeneity between studies (P=0.006, $I^2=80\%$; Figure 3). We also pooled the ORR from five articles with no significant heterogeneity (P=0.41, $I^2=0\%$) and showed the result (OR=1.10, 95% CI=0.89-1.35, P=0.38) using fixed-effect

(0.36–0.89) NA (0.69–1.12) RFS (HR, 95% CI) 0.88 0.57 ₹ ₹ (0.48-0.90) (0.94–1.70) (0.66-0.93) EFS (HR, 95% CI) I.26 0.66 0.78 ₹ ₹ (0.73–I.44) (0.70-1.15) (0.58-1.27) (0.63-0.96) OS (HR, 95% CI) I.03 0.90 0.78 0.86 ₹ **ORR/CG 188/200** 191/357 23/122 78/133 61/95 ORR/EG 277/300 212/360 29/122 58/102 81/134 Generally (years) 18-60 20-81 Age 99 V ≥60 80 AML in first relapse Р Newly diagnosed Patient states Untreated AML secondary AML AML from MDS Untreated AML De novo AML, AML D, C, sorafenib (400 mg twice D, C (7+3 regimen), sorafenib induction, following high-dose Lestaurtinib or not, following -estaurtinib or not, following Midostaurin or placebo, D/C salvage chemotherapy (MEC) nduction and consolidation (400 mg twice daily), or each of four cycles of Table 2 Baseline characteristics of five RCTs included in the meta-analysis (Part 1) daily), or placebo chemotherapy Treatment placebo FLT3 gene mutation nutation Mixed Mixed types FLT3 FLT3 FLT3 Lestaurtinib Lestaurtinib Midostaurin Subgroup Sorafenib Sorafenib Year 2017 2015 2013 2011 2017 ISRNCTNI7161961 SRNCTN55675535 **Clinical register** 2008-004968-40 NCT00079482 NCT00893373 NCT00373373 CALGB10603 RATIFY

Knapper S³⁴

Author

Note: ^aCR was calculated.

NCT00651261

hazard ratio; CI, confidence interval; EFS, event-free survival; RFS, relapse-free survival; AML, acute myeloid leukemia; NA, not available; MEC, mitoxantrone, etoposide, cytarabine; D, daunorubicin; C, cytarabine; MDS, myelodysplastic syndrome; CR, complete remission overall survival; HR, experimental group; CG, control group; OS, overall response rate; EG, Abbreviations: RCT, randomized controlled trial; FLT3, FMS-like tyrosine kinase 3; ORR,

C consolidation

mutation

Röllig C^{36,a}

Levis M³⁵

Serve H³³

Stone RM³⁹

Study or subgroup	log(HR)	SE	Weight	HR IV, Fixed, 95% CI	HR IV, Fixed, 95% CI	
1.1.1 Sorafenib Röllig et al (2015) ³⁶ Serve et al (2013) ³³ Subtotal (95% CI) Heterogeneity: χ^2 =0.46, df=1 (<i>P</i> = Test for overall effect: <i>Z</i> =0.37 (<i>P</i> =0)	-0.1508 0.0296 0.50); <i>I</i> ² =0 ⁶ 0.71)	0.1999 0.1735 %	12.1% 16.0% 28.1%	0.86 (0.58–1.27) 1.03 (0.73–1.45) 0.95 (0.74–1.23)	•	
1.1.2 Lestaurtinib Knapper et al (2017) ³⁴ Subtotal (95% CI) Heterogeneity: Not applicable Test for overall effect: <i>Z</i> =0.83 (<i>P</i> =0	-0.1054 0.41)	0.1267	30.1% 30.1%	0.90 (0.70–1.15) 0.90 (0.70–1.15)	-	
1.1.3 Midostaurin Stone et al (2017) ³⁹ Subtotal (95% CI) Heterogeneity: Not applicable Test for overall effect: <i>Z</i> =2.31 (<i>P</i> =0	-0.2485 0.02)	0.1074	41.8% 41.8%	0.78 (0.63–0.96) 0.78 (0.63–0.96)	*	
Total (95% CI) Heterogeneity: χ^2 =2.04, df=3 (<i>P</i> =0 Test for overall effect: <i>Z</i> =2.15 (<i>P</i> =0 Test for subgroup differences: χ^2 =	0.56); <i>I</i> ²=0% 0.03) 1.57, <i>df</i> =2	% (<i>P</i> =0.46)	100.0% ; / ² =0%	0.86 (0.75–0.99) 0.1 Fa	0.2 0.5 1 2 5 vors FLT3 inhibitors Favors contro	

Figure 2 Forest plot for pooled HR of OS between combined use of FLT3 inhibitors with chemotherapy and chemotherapy alone in patients with AML. Abbreviations: HR, hazard ratio; OS, overall survival; FLT3, FMS-like tyrosine kinase 3; AML, acute myeloid leukemia; SE, standard error; IV, inverse variance; CI, confidence interval.



Figure 3 Forest plot for pooled HR of EFS between combined use of FLT3 inhibitors with chemotherapy and chemotherapy alone in patients with AML. Abbreviations: HR, hazard ratio; EFS, event-free survival; FLT3, FMS-like tyrosine kinase 3; AML, acute myeloid leukemia; SE, standard error; IV, inverse variance; CI, confidence interval.

models (Figure 4). The pooled results showed improved OS with the addition of FLT3 inhibitors to conventional chemotherapy, while no significant improvement on EFS and ORR was seen.

Part I: sensitivity and subgroup analysis

When we changed the fixed-effect models to randomeffect models for pooled analysis of the HR of OS, the results remained the same (HR=0.86, 95% CI=0.75–0.99,

	Chemothera	py+TKI	Chem	otherap	у	OR	OR
Study or subgroup	Events	Total	Events	Total	Weight	M–H, Fixed, 95% CI	M–H, Fixed, 95% CI
1.3.1 Sorafenib Röllig et al (2015) ³⁶	81	134	78	133	18 1%	1 08 (0 66–1 76)	
Serve et al $(2013)^{33}$ Subtotal (95% CI)	58	102 236	61	95 228	15.9% 34.0%	0.73 (0.41–1.30) 0.92 (0.63–1.33)	-
Total events	139		139			· · · · · ·	
Heterogeneity: $\chi^2=0.9$	99, <i>df</i> =1 (<i>P</i> =0	.32): <i>I</i> ² =	=0%				
Test for overall effect	: Z=0.46 (P=0	0.65)					
1.3.2 Lestaurtinib Knapper et al (2017)	34 277	300	188	200	10.1%	0.77 (0.37–1.58)	
Levis et al (2011) ³⁵ Subtotal (95% CI)	29	112 412	23	112 312	9.9% 20.0%	1.35 (0.72–2.52) 1.06 (0.66–1.69)	•
Total events	306		211				
Heterogeneity: $\chi^2=1$.	35, <i>df</i> =1 (<i>P</i> =	0.25): <i>I</i> ²	=26%				
Test for overall effect	: Z=0.24 (P=0).81)					
1.3.3 Midostaurin							+
Stone et al (2017) ³⁹ Subtotal (95% CI)	212	360 360	191	357 357	46.0% 46.0%	1.24 (0.93–1.67) 1.24 (0.93–1.67)	•
Total events	212		191				
Heterogeneity: Not a	oplicable						
Test for overall effect	: Z=1.45 (<i>P</i> =0).15)					•
Total (95% CI)		1008		897	100.0%	1.10 (0.89–1.35)	
Total events	657		541				
Heterogeneity: χ^2 =3.9	95, <i>df</i> =4 (<i>P</i> =0	.41); <i>I</i> ² =	=0%			0.1	Eavors control Eavors ELT3 inhibitors
Test for overall effect	: Z=0.87 (P=0).38)					

Test for subgroup differences: χ^2 =1.62, *df*=2 (*P*=0.44); *I*²=0%

Figure 4 Forest plot for pooled OR of ORR between combined use of FLT3 inhibitors with chemotherapy and chemotherapy alone in patients with AML. Abbreviations: OR, odds ratio; ORR, overall response rate; FLT3, FMS-like tyrosine kinase 3; AML, acute myeloid leukemia; TKI, tyrosine kinase inhibitor; M–H, Mantel–Haenszel; Cl, confidence interval

P=0.03); when we changed to use fixed-effect models for pooled analysis of the HR of EFS, the result showed a significant improvement in EFS in the combined therapy group (HR=0.83, 95% CI=0.73-0.95, P=0.008). Two sorafenib studies included patients with different FLT3 statuses, and a sensitivity analysis on ORR was conducted using the data based on FLT3-ITD patients (data from SORAML study were not changed because no details were extracted), also showing no significant improvement on ORR (OR=1.14, 95% CI=0.92-1.43, P=0.23; heterogeneity P=0.55, I²=0%). The pooled data of subgroup analysis by types of FLT3 inhibitors showed no significant improvement in ORR in the sorafenib subgroup as compared with the current conventional chemotherapy (OR=0.92, 95% CI=0.63-1.33, P=0.65; heterogeneity P=0.32, $I^2=0\%$). In the lestaurtinib subgroup, the pooled ORR (OR=1.06, 95% CI=0.66-1.69, P=0.81; heterogeneity P=0.25, $I^2=26\%$) was also not statistically significant. When we changed the statistical methods to random-effect models or used pool risk ratio (RR), the results were consistent.

Part II: effect of FLT3 inhibitors between FLT3–ITD-positive and -negative AML

A total of 21 studies were included in Part II for studying the different efficacies of FLT3 inhibitors on FLT3–ITD or non-FLT3–ITD AML. We were unable to extract the CR data in one of the articles.¹⁸ In all, 20 studies were pooled to analyze CR and 21 studies were pooled to analyze ORR. In CR analysis, 397 patients with FLT3–ITD and 377 patients without this mutation were included. There was no significant heterogeneity (P=0.97, P=0%). Fixed-effect models were selected, and a forest plot was performed to show the results (OR=1.89, 95% CI=1.06–3.37, P=0.03; Figure 5). The pooled result showed better performance on CR when FLT3 inhibitors were used in FLT3–ITD-positive AML patients compared to FLT3–ITD-negative patients.

Then, we pooled the ORR from all 21 articles. There were 423 patients with FLT3–ITD mutation and 443 patients without this mutation. The heterogeneity (P=0.69, $l^2=0\%$) was not statistically significant, and fixed-effect models were selected. A forest plot was performed to show the results (OR=3.07, 95% CI=2.13–4.43, P<0.001; Figure 6). Our finding showed that FLT3–ITD-positive AML patients were more sensitive to FLT3 inhibitors, benefited more from FLT3 inhibitor treatment and therefore could get better ORR.

Part II: sensitivity and subgroup analysis

Studies about sorafenib, quizartinib, and midostaurin were selected for subgroup analysis. In the sorafenib subgroup,

Study or subgroup	Experimer	ntal Co Total Even	ntrol ts Total	Weight	OR M-H Fixed 95% Cl	OR M-H Fixed 95% CI
4.1.1 Sorafenib Crump et al $(2010)^{15}$ Inaba et al $(2011)^{19}$ Macdonald et al $(201)^{17}$ Pratz et al $(2010)^{11}$ Ravandi et al $(2010)^{12}$ Ravandi et al $(2014)^{2}$ Röllig et al $(2012)^{25}$ Zhang et al $(2008)^{27}$ Subtotal $(95\%$ CI) Total events Heterogeneity: χ^2 =1.0 Test for overall effect:	$ \begin{array}{c} 1 \\ 3 \\ 3)^{21} \\ 0 \\ 4 \\ 12 \\ 3 \\ 17 \\ 2 \\ 1 \\ 36 \\ 18, df=6 (P^{2}) \\ Z=2.49 (P^{2}) \end{array} $	9 5 1 2 13 2 19 3 22 7 7 78 6 =0.98); <i>I</i> ² =0 =0.01)	0 18 3 6 1 20 0 10 6 38 2 43 0 6 0 9 150 2 %	1.6% 6.0% 1.1% 5.6% 11.4% 3.8% 2.0% 31.5%	6.53 (0.24–177.39) 1.50 (0.14–16.54) 4.33 (0.12–159.52) Not estimable 5.54 (0.64–47.62) 2.92 (0.58–14.73) 1.59 (0.07–37.44) 4.38 (0.15–125.29) 3.28 (1.29–8.37)	
4.1.2 Quizartinib Cooper et al (2016) ¹³ Cortes et al (2012) ³⁰ Cortes et al (2013) ¹⁴ Levis et al (2012) ²⁹ Subtotal (95% CI) Total events Heterogeneity: χ^2 =2.8 Test for overall effect:	1 0 1 4 6 1, <i>df</i> =3 (<i>P</i> : <i>Z</i> =0.23 (<i>P</i>)	7 92 17 99 215 =0.42); / ² =0 =0.82)	1 7 1 41 0 37 1 38 123 3 %	4.7% 11.4% 1.6% 7.7% 25.4%	1.00 (0.05–19.96) 0.15 (0.01–3.66) 6.82 (0.26–176.29) 1.56 (0.17–14.40) 1.16 (0.33–4.07)	
4.1.3 Midostaurin Ramsingh et al $(2014)^{26}$ Valker et al $(2012)^{26}$ Walker et al $(2016)^{28}$ Subtotal (95% CI) Total events Heterogeneity: $\chi^2=0.7$ Total for overall effect) ²² 1 8 3 12 '2, df=2 (P: : Z=0.19 (F	5 9 2 6 1 20 =0.70); <i>I</i> ² =0 2=0.85)	1 4 4 31 0 17 52 5 %	4.9% 6.6% 14.4% 26.0%	0.75 (0.03–17.51) 2.33 (0.25–21.98) 0.70 (0.11–4.54) 1.13 (0.32–3.93)	
4.1.4 Sunitinib Fiedler et al (2005) ¹⁷ Fiedler et al (2015) ¹⁶ Subtotal (95% CI) Total events Heterogeneity: Not ap Total for overall effect	0 7 pplicable : <i>Z</i> =0.56 (<i>F</i>	2 14 16 P=0.57)	0 10 3 8 18 3	10.6% 10.6%	Not estimable 1.67 (0.28–9.82) 1.67 (0.28–9.82)	
4.1.5 Lestaurtinib Knapper et al (2006) ² Subtotal (95% CI) Total events Heterogeneity: Not ap Total for overall effect	⁰ 0 0 pplicable : Not applic	2 2 cable	0 25 25 0		Not estimable Not estimable	
4.1.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not ap Total for overall effect	1 1 pplicable : Z=0.60 (F	2 2 P=0.55)	1 4 4 1	1.8% 1.8%	3.00 (0.08–107.45) 3.00 (0.08–107.45)	,
4.1.7 Multiple types Alvarado et al (2014) Subtotal (95% CI) Total events Heterogeneity: Not ap Total for overall effect	¹² 4 4 oplicable : Z=0.13 (F	64 64 P=0.90)	0 5 5 0	4.7% 4.7%	0.82 (0.04–17.26) 0.82 (0.04–17.26)	
Total (95% CI) Total events Heterogeneity: χ^2 =7.3 Total for overall effect Test for subgroup diff	66 32, <i>df</i> =16 (<i>f</i> : Z=2.17 (<i>F</i> erences: χ ²	397 =0.97); <i>I</i> ² =0 =0.03) = 2.94, df=	377 4)% 5 (<i>P</i> =0.7	100.0% 71); <i>I</i> ² =0%	1.89 (1.06–3.37) – – – – – – – – – – – – – – – – – – –	0.1 1 10 100 Favors control Favors FLT3 inhibitors

Figure 5 Forest plot for pooled OR of CR between FLT3–ITD-positive patients and FLT3–ITD-negative patients. Abbreviations: OR, odds ratio; CR, complete remission; FLT3, FMS-like tyrosine kinase 3; ITD, internal tandem duplication; M–H, Mantel–Haenszel; CI, confidence interval.

	Experime	ntal	Contro	bl		OR	OR
Study or subgroup	Events	Total	Events	Total	Weight	M–H, Fixed, 95% CI	M–H, Fixed, 95% Cl
4.2.1 Sorafenib							
Crump et al (2010) ¹⁵	1	9	0	18	0.9%	6.53 (0.24, 177.39)	
Inaba et al (2011) ¹⁹	5	5	4	6	1.0%	6.11 (0.23, 162.73)	
Macdonald et al (2013)	²¹ 1	1	1	20	0.2%	39.00 (1.06, 1435.65)	
Pratz et al (2010) ¹¹	0	2	0	10		Not estimable	
Ravandi et al (2010) ²⁴	13	13	29	38	1.6%	8.69 (0.47, 160.54)	
Ravandi et al (2014) ²³	18	19	36	43	3.4%	3.50 (0.40, 30.66)	
Röllig et al (2012) ²⁵	5	22	1	6	3.5%	1.47 (0.14, 15.69)	
Zhang et al (2008) ²⁷	6	7	3	9	1.1%	12.00 (0.96, 150.69)	
Subtotal (95% CI)		78		150	11.7%	5.42 (1.91, 15.41)	
Total events	49		74			(, , , , , , , , , , , , , , , , , , ,	
Heterogeneity: $\chi^2 = 2.97$, df=6 (P=0).81): <i>I</i> ²:	=0%				
Test for overall effect: Z	=3.17 (<i>P</i> =0	0.001)					
4.2.2 Quizartinib							
Cooper et al (2016) ¹³	3	7	1	7	1.7%	4.50 (0.34, 60.15)	
Cortes et al (2012) ³⁰	66	92	17	41	19.4%	3.58 (1.66, 7.74)	
Cortes et al (2013) ¹⁴	9	17	5	37	4.3%	7.20 (1.89, 27.49)	
Levis et al (2012) ²⁹	67	99	18	38	24.5%	2.33 (1.08, 4.99)	
Subtotal (95% CI)		215		123	49.9%	3.31 (2.03, 5.40)	•
Total events	145		41			,	
Heterogeneity: $\gamma^2 = 2.21$. df=3 (P=0).53): / ² =	=0%				
Test for overall effect: Z	=4.79 (P<(0.00001)				
			,				
4.2.3 Midostaurin							
Fischer et al (2010) ¹⁸	19	26	30	66	13.3%	3.26 (1.21, 8.79)	
Ramsingh et al (2014) ²	² 2	5	1	4	1.9%	2 00 (0 11 35 81)	
Stone et al (2012) ²⁶	8	9	24	31	3.5%	2 33 (0 25 21 98)	
Walker et al (2016) ²⁸	4	6	15	17	7.6%	0.27 (0.03, 2.53)	
Subtotal (95% CI)		46	10	118	26.4%	2 18 (0 99 4 77)	
Total overte	33	40	70		20.4 /0	2.10 (0.33, 4.77)	
Heterogeneity: x ² =3.00	df=3 (P=0	26) · 12-	-25%				
Test for overall effect: Z	=1.95 (<i>P</i> =0	0.05)	2070				
4.2.4 Sunitinib							
Fiedler et al (2005) ¹⁷	2	2	3	10	0.7%	10.71 (0.40, 287.83)	
Fiedler et al (2015) ¹⁶	9	14	6	8	8.0%	0.60 (0.09 4.17)	
Subtotal (95% CI)	Ŭ	16	0	18	8.7%	1 45 (0 34 6 22)	
Total events	11	10	9		0.1 /0	1.10 (0.01, 0.22)	
Heterogeneity: $\gamma^2 = 2.22$	df=1 (P=0) 14)· <i>I</i> ² :	-55%				
) 62)	00,0				
Test for overall effect: Z	=0.50 (P=0	,)					
Test for overall effect: Z	=0.50 (<i>P</i> =0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,					
Test for overall effect: Z 4.2.5 Lestaurtinib	=0.50 (<i>P</i> =0	,,					
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰	(P=0.50 (P=0)	2	0	25		Not estimable	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI)	(=0.50 (<i>P</i> =0)	2 2	0	25 25		Not estimable Not estimable	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events	(P=0.50 (P=0) 0 0	2 2	0	25 25		Not estimable Not estimable	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app	=0.50 (<i>P</i> =(0 licable	2 2	0 0	25 25		Not estimable Not estimable	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N	=0.50 (<i>P</i> =(0 licable lot applicat	2 2 2	0	25 25		Not estimable Not estimable	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N	=0.50 (<i>P</i> =(0 licable lot applicat	2 2 2	0 0	25 25		Not estimable Not estimable	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib	=0.50 (<i>P</i> =(0 licable 'ot applicat	2 2 ble	0	25 25		Not estimable Not estimable	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% Cl) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹	=0.50 (<i>P</i> =(0 licable lot applicat	2 2 Dile 2	0 0 3	25 25 4	1.3%	Not estimable Not estimable 2.14 (0.06, 77.54)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI)	=0.50 (<i>P</i> =(0 licable lot applicat	2 2 ble 2 2	0 0 3	25 25 4 4	1.3% 1.3%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events	e=0.50 (P=(0 licable lot applicat	2 2 ble 2 2	0 0 3 3	25 25 4 4	1.3% 1.3%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib Iyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app	e=0.50 (P=(0 licable lot applicat 2 licable	2 2 ble 2 2	0 0 3 3	25 25 4 4	1.3% 1.3%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 licable =0.42 (<i>P</i> =(2 2 ble 2 2	0 0 3 3	25 25 4 4	1.3% 1.3%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib Iyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z	=0.50 (<i>P</i> =(0 licable lot applicat 2 licable =0.42 (<i>P</i> =(2 2 ble 2 2 2	0 0 3 3	25 25 4 4	1.3% 1.3%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types	=0.50 (<i>P</i> =(0 licable lot applicat 2 licable =0.42 (<i>P</i> =(2 2 ble 2 2 0.68)	0 0 3 3	25 25 4 4	1.3% 1.3%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types Alvarado et al (2014) ¹²	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 2 licable =0.42 (<i>P</i> =(13	2 2 2 2 2 2 2 2 2 0.68) 64	0 0 3 3 0	25 25 4 4 5	1.3% 1.3% 2.1%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types Alvarado et al (2014) ¹² Subtotal (95% CI)	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 licable =0.42 (<i>P</i> =(13	2 2 2 0le 2 2 0.68) 64 64	0 0 3 3 0	25 25 4 4 5 5	1.3% 1.3% 2.1% 2.1%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45)	
Test for overall effect: <i>Z</i> 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: <i>Z</i> 4.2.7 Multiple types Alvarado et al (2014) ¹² Subtotal (95% CI) Total events	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 2 licable =0.42 (<i>P</i> =(13 13	2 2 2 0le 2 2 0.68) 64 64	0 0 3 3 0 0	25 25 4 4 5 5	1.3% 1.3% 2.1% 2.1%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib Iyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types Alvarado et al (2014) ¹² Subtotal (95% CI) Total events Heterogeneity: Not app	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 licable =0.42 (<i>P</i> =(13 13 licable	2 2 2 0le 2 2 0.68) 64 64	0 0 3 3 3 0 0	25 25 4 4 5 5	1.3% 1.3% 2.1% 2.1%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% Cl) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% Cl) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types Alvarado et al (2014) ¹² Subtotal (95% Cl) Total events Heterogeneity: Not app Test for overall effect: Z	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 2 licable =0.42 (<i>P</i> =(13 13 licable =0.70 (<i>P</i> =(2 2 ble 2 2 0.68) 64 64 0.48)	0 0 3 3 0 0	25 25 4 4 4 5 5	1.3% 1.3% 2.1% 2.1%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types Alvarado et al (2014) ¹² Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z Subtotal (95% CI)	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 2 licable =0.42 (<i>P</i> =(13 13 licable =0.70 (<i>P</i> =(2 2 ble 2 2 0.68) 64 64 0.48) 2.48)	0 0 3 3 3 0 0	25 25 4 4 4 4 5 5	1.3% 1.3% 2.1% 2.1%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib lyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types Alvarado et al (2014) ¹² Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z Utal events Heterogeneity: Not app Test for overall effect: Z	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 2 licable =0.42 (<i>P</i> =(13 13 licable =0.70 (<i>P</i> =(253	2 2 2 0.68) 64 64 0.48) 423	0 0 3 3 3 0 0	25 25 4 4 4 5 5 5	1.3% 1.3% 2.1% 2.1%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45) 3.07 (2.13, 4.43)	
Test for overall effect: Z 4.2.5 Lestaurtinib Knapper et al (2006) ²⁰ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: N 4.2.6 Crenolanib Iyer et al (2016) ³¹ Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z 4.2.7 Multiple types Alvarado et al (2014) ¹² Subtotal (95% CI) Total events Heterogeneity: Not app Test for overall effect: Z Total (95% CI) Total events	=0.50 (<i>P</i> =(0 0 licable lot applicat 2 2 licable =0.42 (<i>P</i> =(13 13 licable =0.70 (<i>P</i> =(253 8, df=19 (<i>T</i>)	2 2 ble 2 2 0.68) 64 64 0.48) 423 -0.60	0 0 3 3 0 0 0	25 25 4 4 4 5 5 443	1.3% 1.3% 2.1% 2.1% 100.0%	Not estimable Not estimable 2.14 (0.06, 77.54) 2.14 (0.06, 77.54) 2.88 (0.15, 55.45) 2.88 (0.15, 55.45) 3.07 (2.13, 4.43)	

Figure 6 Forest plot for pooled OR of ORR between FLT3–ITD-positive patients and FLT3–ITD-negative patients. Abbreviations: OR, odds ratio; ORR, overall response rate; FLT3, FMS-like tyrosine kinase 3; ITD, internal tandem duplication; M–H, Mantel–Haenszel; CI, confidence interval. eight relevant studies were pooled, showing better CR (OR=3.28, 95% CI=1.29-8.37, P=0.01; heterogeneity P=0.98, $I^2=0\%$) and ORR (OR=5.42, 95% CI=1.91-15.41, P=0.001; heterogeneity P=0.81, $I^2=0\%$) in patients with FLT3-ITD-positive AML. In the quizartinib subgroup, four relevant studies were pooled, showing that patients with FLT3-ITD-positive AML got better ORR (OR=3.31, 95% CI=2.03-5.40, P < 0.001; heterogeneity P = 0.53, $I^2 = 0\%$). Quizartinib did not seem to improve CR significantly (OR=1.16, 95% CI=0.33-4.07, P=0.82; heterogeneity P=0.42, I²=0%) in FLT3-ITD-positive AML patients. No significant preference of CR (OR=1.13, 95% CI=0.32-3.93, P=0.85; heterogeneity P=0.70, $I^2=0\%$) and ORR (OR=2.18, 95% CI=0.99-4.77, P=0.05; heterogeneity P=0.26, I²=25%) in FLT3-ITD-positive AML was observed in the midostaurin subgroup (Figures 5 and 6). Subgroup analysis was also carried out based on primary or relapsed AML. In the primary AML subgroup, six relevant studies were pooled, showing better CR (OR=2.90, 95% CI=1.17-7.18, P=0.02; heterogeneity P=0.94, I²=0%) and ORR (OR=2.78, 95%) CI=1.06-7.32, P=0.04; heterogeneity P=0.28, I²=22%) in FLT3-ITD-positive patients. In the relapsed AML subgroup, 15 studies were included and FLT3-ITD-positive patients had better ORR (OR=3.13, 95% CI=2.11-4.65, P<0.001; heterogeneity P=0.74, $I^2=0\%$), whereas no significant preference of CR (OR=1.34, 95% CI=0.63-2.88, P=0.45; heterogeneity P=0.93, $I^2=0\%$) was detected. In all, 14 studies with >20 participants included were selected for sensitivity analysis; the results remained the same in either CR (OR=1.96, 95% CI=1.03-3.72, P=0.04; heterogeneity P=0.84, $I^2=0\%$) or ORR (OR=2.88, 95% CI=1.96-4.24, P<0.001; heterogeneity P=0.41, P=4%).

Publication bias

Because only five studies were included in Part I, funnel plots had low power and were not used in test for publication bias. In Part II, we obtained symmetric funnel plots for the synthesis of CR and ORR, indicating mild publication bias (Figure S2A and B).

Discussion

AML is a hematologic malignancy characterized by multiple cytogenetic and genetic abnormalities in the hematopoietic stem or progenitor cells.⁵ As a common mutation, FLT3–ITD was detected in 20%–30% of AML patients and reported to be associated with worse prognosis.⁴⁰ Patients with FLT3–ITD mutations and a normal karyotype are considered in the poorrisk group according to the National Comprehensive Cancer

Network (NCCN) guidelines. In the genetic risk classification of the 2017 European LeukemiaNet (ELN) recommendation, a high FLT3–ITD allelic ratio is thought to be associated with a high relapse rate and poor OS. Patients with NPM1 mutation and FLT3–ITD with a low allelic ratio belong to the favorable risk group, while AML patients with wild-type NPM1 and FLT3–ITD with a high allelic ratio have a poor prognosis and are placed in the adverse-risk group.^{41,42} FLT3 inhibitors have been applied to target mutant FLT3 and block related pathways potentially involved in the development and progression of malignancies.⁴³ However, there is no consensus on the use of FLT3 inhibitors, and most of these compounds are "off-label" used in AML.

FLT3 inhibitors undergoing clinical evaluation were divided into the first-generation agents (including sunitinib, midostaurin, and lestaurtinib) with relative nonspecificity and multiple targets and the second-generation agents (including sorafenib, quizartinib, and crenolanib) with a higher selectivity against mutant FLT3.32 A host of first-generation FLT3 inhibitors were limited by their suboptimal propensity for AML control in vivo or their off-target effects, frequently producing a peripheral blast cell clearance and BM blast cell persistence.^{10,44} In our study, no patient was found to achieve ORR with lestaurtinib monotherapy and 41.7% patients responded to sunitinib monotherapy with the best outcome of PR.17,20 It was documented that midostaurin as a single agent could reduce blast count in peripheral blood in 70% patients, and the peripheral blood and BM BR was shown in 53.3% patients with monotherapy in studies from our analysis.^{18,45} For second-generation FLT3 inhibitors, quizartinib monotherapy has been reported with composite CR (CRc; including CR, CRi, and CRp) in 46-57% patients with relapsed/refractory (R/R) AML.^{14,46,47} Crenolanib, as a single agent, achieved the CRc up to 39% in this patient population.^{48–50} The median OS for quizartinib and crenolanib in initial clinical studies was reported up to 168 and 259 days, and median EFS could achieve 74 and 56 days, respectively, which is similar or better than conventional chemotherapy with median OS of 198 days and 4-month EFS ranging from 16.6% to 37.7%.50-52 Improvement on EFS and RFS has also been seen from the use of sorafenib.³⁶ In the studies included in our analysis, >50% the patients using sorafenib and quizartinib achieved ORR, and CR in two subgroups was 43.0% and 2.66%, respectively, in FLT3-ITD-positive and -negative patients. The reason of much higher CR in the sorafenib subgroup might be that 75.9% (173/228) patients in the sorafenib subgroup received combined therapy regimen, whereas 95.9% (324/338) patients used quizartinib as a

single agent. These results demonstrate notable and promising clinical efficacy of second-generation FLT3 inhibitors.

Compared with conventional chemotherapy, addition of FLT3 inhibitors seems to improve the OS of AML patients, although no significantly improved EFS and ORR were detected. When subgroup analysis was carried out on different FLT3 inhibitors, midostaurin was the only agent with a significantly higher OS and EFS, based on the results of CALGB 10603/RATIFY study.37 Midostaurin became the first FLT3 inhibitor for the treatment of FLT3-mutant AML, approved by the FDA on April 28, 2017 and the European Medicines Agency (EMA) on July 20, 2017. The results of the SORAML study³⁶ showed that EFS and RFS were prolonged when sorafenib was added to standard chemotherapy in patients aged 60 years or younger with newly diagnosed AML, suggesting that sorafenib might have the potential to reduce relapses and increase the time in CR. However, another sorafenib study by Serve et al³³ reported different results, which might be attributed to older age (>60 years) of the participants. Considering that FLT3-ITD-positive patients might be more sensitive to these agents, the benefit of sorafenib on AML might be underestimated in these two studies, because both FLT3-mutated and FLT3-WT patients were included and FLT3-ITD-positive patients were a small part of all participants. No improved ORR was found in the pooled data or in each subgroup. It was documented that ~50-80% primary AML patients and 18-60% R/R AML patients could respond to conventional chemotherapy.53 The pooled ORR of combined use of sorafenib, lestaurtinib, and midostaurin in our analysis was 58.9%, 74.3%, and 58.9%, respectively, which might be influenced by the AML status of participants, and these results were consistent with the data of previous conventional chemotherapy. From the results of our analysis, FLT3 inhibitors seem to improve the OS of AML.

FLT3–ITD-positive patients might be more sensitive to FLT3 inhibitors with a higher CR and ORR compared to patients with other mutations or wild type when FLT3 inhibitors were used as combined therapy or monotherapy. This result could be explained by the block of FLT3–ITD-related oncogenic targets and signaling, leading to the suppression of leukemia blast cells.¹⁰ It has been documented that with conventional chemotherapy regimens, FLT3–ITD leads to a decreased disease-free survival (DFS) and OS, although the CR is comparable to that achieved in those with FLT3– WT.^{24,43} To clarify the potential long-term benefit of FLT3 inhibitors between FLT3–ITD-positive and -negative patients, the end points OS and DFS were reviewed and summarized from seven included studies (including three on midostaurin,

three on quizartinib and one on sorafenib) where survival data on different gene types could be extracted.^{14,18,22,23,26,29,30} However, the quantitative analysis is inappropriate due to their various survival data forms. A Phase IIB trial of midostaurin¹⁸ showed similar results to conventional chemotherapy, where the mutant population was with a median OS of 100 days (95% CI=76-121 days), shorter than that in wild-type population of 159 days (95% CI=130-209 days), and it was the only study that provided statistical difference. A Phase II sorafenib clinical trial²³ showed that the median OS and DFS for patients with FLT3-ITD was 15.5 and 9.9 months, respectively, shorter than for those without FLT3-ITD. Additional two studies on midostaurin reported the similar result that DFS and OS probabilities in patients with mutation were similar to those with FLT3-WT disease.^{22,26} Inconsistent results that whether FLT3-ITD-positive or -negative patients would have better OS and DFS were shown in quizartinib.14,29,30 For crenolanib, Cortes et al48 and Randhawa et al⁴⁹ reported that the OS in mutant patients was longer than that in patients with negative FLT3-ITD, which might be due to the possible increased likelihood of patients able to undergo an allogeneic HCT.36 In contrast to previous chemotherapy reporting significantly worse survival for patients with FLT3-ITD, use of FLT3 inhibitors seems to improve their long-term outcome, although there is no definitive evidence showing that their survival can be equal to or even better than those with negative FLT3-ITD.

Although our results showed that FLT3 inhibitors offered favorable outcomes in FLT3-ITD-positive patients, we cannot conclude that patients with negative FLT3-ITD would not benefit from the use of these compounds. Many initial clinical trials of FLT3 inhibitors were carried out in a mixed population, including FLT3-ITD, FLT3-TKD, and FLT3-WT patients, which allowed the investigators to get safety data and establish whether there existed some signaling pathway to justify the use of FLT3 inhibitors in non-FLT3-mutated patients. In some studies, FLT3-WT AML patients were reported to benefit from the use of these drugs and achieve clinical response when FLT3 inhibitors were used as monotherapy.^{2,18,29} Most FLT3 inhibitors are multi-kinase inhibitors functionally and have different targets. Although FLT3 has been shown to be a key mediator of leukemic cell growth and survival, a great number of related and parallel signal transduction pathways contribute to leukemic progression, many of which are likely attenuated by FLT3 inhibitors.44 However, the relative nonspecificity may also contribute to the adverse effects associated with these drugs, which may limit their clinical utility.

Besides the FLT3 status, several other factors might impact the outcome of AML patients with FLT3 inhibitor treatment. Combined use of FLT3 inhibitors with chemotherapy might improve the clinical outcome, which is also seen in the sorafenib subgroup. From five studies^{11,15,19,25,27} focusing sorafenib on R/R AML, the ORR was 18.2% and 38.5% in monotherapy and combined therapy, respectively. All the completed large-size RCTs³³⁻³⁷ added FLT3 inhibitors to conventional chemotherapy, and combined use of sorafenib with chemotherapy in patients with R/R AML disease was recommended from the NCCN guidelines. Both the pharmacological action and the better physical status in those who can tolerate chemotherapy might contribute to this fact. It has been reported that FLT3 inhibitors are less effective in the relapsed patients compared with the newly diagnosed patients, which is also seen in our analysis.53,54 In Part I, both lestaurtinib studies focused on the addition of lestaurtinib in FLT3-mutant AML patients. The ORR was higher in primary AML than that in relapsed AML (92.3% vs 23.8% in the lestaurtinib group; 94.0% vs 18.9% in the control group). In Part II, six out of 21 studies included only primary AML, and patients from these six studies had higher CR (58.3% vs 7.26%) and ORR (65.0% vs 47.4%). The dramatic increase in the FLT3 ligand level after chemotherapy represents a possible obstacle to inhibiting FLT3.54 Prior therapies might be a key predictor for R/R AML. Ohanian et al⁵⁰ reported that using crenolanib as a single agent, a much longer median OS was observed in patients with ≤ 2 prior therapies (259 days) than in those with ≥ 3 prior therapies (53 days). Randhawa et al⁴⁹ also found that patients without prior FLT3 inhibitor therapy had better CRi, OS, and EFS. Current studies seem to suggest that old patients, especially those older than 60 years, may have worse prognosis. Two RCTs^{33,36} reported the opposite conclusion with sorafenib, saying that patients ≤ 60 years benefitted from the use of sorafenib with improved EFS and RFS. For R/R AML using crenolanib as a single agent, patients <60 years had a higher OS than those \geq 60 years (234 vs 185 days).48 It has been documented that AML in the elderly is genetically heterogeneous and often follows an MDS. However, Perl et al⁵⁵ reported that patients aged 70 years with chemotherapy-resistant AML also preserved high response rates and promising survival to quizartinib.

FLT3 inhibitor therapy is associated with an incidence of several adverse effects in clinical trials. Although various drug-related events have been detected, including gastrointestinal symptoms, cardiac toxicity, infections, fatigue, peripheral edema, mucositis, depression, neuropathy, arthralgia, and skin toxicity,^{15,21} FLT3 inhibitors were generally safe and well tolerated in these Phase I or II clinical trials.^{17,18,20,23,26} The most frequent treatment-related events were grades 1 and 2, which were manageable through treatment interruptions, dose reductions, and medical intervention.¹⁴ Nevertheless, compared with placebo, FLT3 inhibitors might increase the incidence of adverse effects when they were combined with conventional chemotherapy. Two RCTs showed that the addition of sorafenib might lead to more premature termination of study treatment and less consolidation chemotherapy.^{33,36} Similar results were seen in lestaurtinib; the AML15 study showed that there were moderate increases in nausea and diarrhea with the addition of lestaurtinib treatment.³⁴ Levis et al³⁵ reported that 24% of patients receiving combined lestaurtinib therapy discontinued their planned therapy before completion because of adverse events. While for midostaurin, Stone et al³⁷ found that there was no significant difference on grades 3 or 4 therapy-related adverse effects compared with placebo. Therefore, high attention should be paid to the treatmentrelated events when FLT3 inhibitors are used in AML.

Several limitations in our analysis should be mentioned. First, only five studies with three types of FLT3 inhibitors were included in Part I. Although the heterogeneity is acceptable, the agent types, age of participants, either monotherapy or combined therapy and AML status including no prior treatment or relapse are not the same between the studies. Therefore, clinical decisions should be made by taking into account multiple factors. Second, the definition of EFS is similar in the studies of Stone et al³⁷ and Röllig et al³⁶ as primary treatment failure or relapse or death. However, we could not find the exact definition of EFS from the study of Serve et al.³³ This may bring potential bias to our study. Third, due to limited patients included and various responses to different FLT3 inhibitor therapies, results of these studies were inconclusive, leading to relatively large CIs of our pooled analysis. More high-quality studies are needed to validate our conclusions. Finally, we pooled ORR including CR, CRi, and CRp in Part I, and pooled ORR also included PR in Part II. This difference makes it inappropriate to make direct comparison between the data in Part I and those in Part II. Most of the 21 studies in Part II lacked long-term survival data such as OS, EFS, and RFS, and therefore, it is inappropriate to pool them together.

Conclusion

In this meta-analysis, we showed that addition of FLT3 inhibitors to conventional chemotherapy seems to improve the OS of AML patients. FLT3–ITD-positive patients might be more sensitive to FLT3 inhibitors with a higher CR and ORR. FLT3 inhibitors could still give hope to patients who are not candidates for conventional chemotherapy. Based on these findings, we recommend the initiation of more studies to confirm the value of FLT3 inhibitors in AML therapy.

Author contributions

All authors contributed to the data analysis and drafting and critically revising the paper, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure

The authors report no conflicts of interest in this work.

References

- Percival ME, Lai C, Estey E, Hourigan CS. Bone marrow evaluation for diagnosis and monitoring of acute myeloid leukemia. *Blood Rev.* 2017;31(4):185–192.
- 2. Yu MG, Zheng HY. Acute myeloid leukemia: advancements in diagnosis and treatment. *Chin Med J (Engl)*. 2017;130(2):211–218.
- 3. Smith CC, Wang Q, Chin CS, et al. Validation of ITD mutations in FLT3 as a therapeutic target in human acute myeloid leukaemia. *Nature*. 2012;485(7397):260–263.
- Rebechi MT, Pratz KW. Genomic instability is a principle pathologic feature of FLT3 ITD kinase activity in acute myeloid leukemia leading to clonal evolution and disease progression. *Leuk Lymphoma*. 2017;58(9):1–11.
- 5. Estey EH. Acute myeloid leukemia: 2013 update on risk-stratification and management. *Am J Hematol.* 2013;88(4):318–327.
- Port M, Bottcher M, Thol F, et al. Prognostic significance of FLT3 internal tandem duplication, nucleophosmin 1, and CEBPA gene mutations for acute myeloid leukemia patients with normal karyotype and younger than 60 years: a systematic review and meta-analysis. *Ann Hematol.* 2014;93(8):1279–1286.
- Azari-Yam A, Tavakkoly-Bazzaz J, Semnani Y, et al. FLT3 gene mutation profile and prognosis in adult acute myeloid leukemia. *Clin Lab.* 2016;62(10):2011–2017.
- Hassanein M, Almahayni MH, Ahmed SO, Gaballa S, El FR. FLT3 inhibitors for treating acute myeloid leukemia. *Clin Lymphoma Myeloma Leuk*. 2016;16(10):543–549.
- Hu S, Niu H, Minkin P, et al. Comparison of antitumor effects of multitargeted tyrosine kinase inhibitors in acute myelogenous leukemia. *Mol Cancer Ther.* 2008;7(5):1110–1120.
- Hospital MA, Green AS, Maciel TT, et al. FLT3 inhibitors: clinical potential in acute myeloid leukemia. *Onco Targets Ther.* 2017;10: 607–615.
- Pratz KW, Cho E, Levis MJ, et al. A pharmacodynamic study of sorafenib in patients with relapsed and refractory acute leukemias. *Leukemia*. 2010;24(8):1437–1444.
- 12. Alvarado Y, Kantarjian HM, Luthra R, et al. Treatment with FLT3 inhibitor in patients with FLT3-mutated acute myeloid leukemia is associated with development of secondary FLT3-tyrosine kinase domain mutations. *Cancer*. 2014;120(14):2142–2149.
- Cooper TM, Cassar J, Eckroth E, et al. A phase I study of quizartinib combined with chemotherapy in relapsed childhood leukemia: a therapeutic advances in childhood leukemia & lymphoma (TACL) study. *Clin Cancer Res.* 2016;22(16):4014–4022.
- 14. Cortes JE, Kantarjian H, Foran JM, et al. Phase I study of quizartinib administered daily to patients with relapsed or refractory acute myeloid leukemia irrespective of FMS-like tyrosine kinase 3-internal tandem duplication status. *J Clin Oncol.* 2013;31(29):3681–3687.

- Crump M, Hedley D, Kamel-Reid S, et al. A randomized phase I clinical and biologic study of two schedules of sorafenib in patients with myelodysplastic syndrome or acute myeloid leukemia: a NCIC (National Cancer Institute of Canada) Clinical Trials Group Study. *Leuk Lymphoma*. 2010;51(2):252–260.
- Fiedler W, Kayser S, Kebenko M, et al. A phase I/II study of sunitinib and intensive chemotherapy in patients over 60 years of age with acute myeloid leukaemia and activating FLT3 mutations. *Br J Haematol.* 2015;169(5):694–700.
- Fiedler W, Serve H, Dohner H, et al. A phase 1 study of SU11248 in the treatment of patients with refractory or resistant acute myeloid leukemia (AML) or not amenable to conventional therapy for the disease. *Blood*. 2005;105(3):986–993.
- Fischer T, Stone RM, Deangelo DJ, et al. Phase IIB trial of oral midostaurin (PKC412), the FMS-like tyrosine kinase 3 receptor (FLT3) and multi-targeted kinase inhibitor, in patients with acute myeloid leukemia and high-risk myelodysplastic syndrome with either wild-type or mutated FLT3. *J Clin Oncol.* 2010;28(28):4339–4345.
- Inaba H, Rubnitz JE, Coustan-Smith E, et al. Phase I pharmacokinetic and pharmacodynamic study of the multikinase inhibitor sorafenib in combination with clofarabine and cytarabine in pediatric relapsed/ refractory leukemia. *J Clin Oncol.* 2011;29(24):3293–3300.
- Knapper S, Burnett AK, Littlewood T, et al. A phase 2 trial of the FLT3 inhibitor lestaurtinib (CEP701) as first-line treatment for older patients with acute myeloid leukemia not considered fit for intensive chemotherapy. *Blood*. 2006;108(10):3262–3270.
- Macdonald DA, Assouline SE, Brandwein J, et al. A phase I/II study of sorafenib in combination with low dose cytarabine in elderly patients with acute myeloid leukemia or high-risk myelodysplastic syndrome from the National Cancer Institute of Canada Clinical Trials Group: trial IND186. *Leuk Lymphoma*. 2013;54(4):760–766.
- 22. Ramsingh G, Westervelt P, McBride A, et al. Phase I study of cladribine, cytarabine, granulocyte colony stimulating factor (CLAG regimen) and midostaurin and all-trans retinoic acid in relapsed/refractory AML. *Int J Hematol.* 2014;99(3):272–278.
- Ravandi F, Arana YC, Cortes JE, et al. Final report of phase II study of sorafenib, cytarabine and idarubicin for initial therapy in younger patients with acute myeloid leukemia. *Leukemia*. 2014;28(7):1543–1545.
- Ravandi F, Cortes JE, Jones D, et al. Phase I/II study of combination therapy with sorafenib, idarubicin, and cytarabine in younger patients with acute myeloid leukemia. *J Clin Oncol.* 2010;28(11):1856–1862.
- Röllig C, Brandts C, Shaid S, et al. Survey and analysis of the efficacy and prescription pattern of sorafenib in patients with acute myeloid leukemia. *Leuk Lymphoma*. 2012;53(6):1062–1067.
- 26. Stone RM, Fischer T, Paquette R, et al. Phase IB study of the FLT3 kinase inhibitor midostaurin with chemotherapy in younger newly diagnosed adult patients with acute myeloid leukemia. *Leukemia*. 2012;26(9):2061–2068.
- Zhang W, Konopleva M, Shi YX, et al. Mutant FLT3: a direct target of sorafenib in acute myelogenous leukemia. *J Natl Cancer Inst.* 2008;100(3):184–198.
- Walker AR, Wang H, Walsh K, et al. Midostaurin, bortezomib and MEC in relapsed/refractory acute myeloid leukemia. *Leuk Lymphoma*. 2016;57(9):2100–2108.
- 29. Levis MJ, Perl AE, Dombret H, et al. Final results of a phase 2 openlabel, monotherapy efficacy and safety study of quizartinib (AC220) in patients with FLT3-ITD positive or negative relapsed/refractory acute myeloid leukemia after second-line chemotherapy or hematopoietic stem cell transplantation. *Blood*. 2012;120(21):673.
- 30. Cortes JE, Perl AE, Dombret H, et al. Final results of a phase 2 openlabel, monotherapy efficacy and safety study of quizartinib (AC220) in patients 60 years of age with FLT3 ITD positive or negative relapsed/ refractory acute myeloid leukemia. *Blood*. 2012;120(21):48.
- Iyer SP, Jethava Y, Karanes C, Eckardt JR, Collins R. Safety study of salvage chemotherapy high-dose ara-C/mitoxantrone (HAM) and type I FLT3-TKI crenolanib in first relapsed/primary refractory AML. *Blood*. 2016;128(22):3983–3983.

- Fathi AT, Chen YB. The role of FLT3 inhibitors in the treatment of FLT3mutated acute myeloid leukemia. *Eur J Haematol*. 2017;98(4):330–336.
- 33. Serve H, Krug U, Wagner R, et al. Sorafenib in combination with intensive chemotherapy in elderly patients with acute myeloid leukemia: results from a randomized, placebo-controlled trial. *J Clin Oncol.* 2013;31(25):3110–3118.
- Knapper S, Russell N, Gilkes A, et al. A randomized assessment of adding the kinase inhibitor lestaurtinib to first-line chemotherapy for FLT3-mutated AML. *Blood*. 2017;129(9):1143–1154.
- 35. Levis M, Ravandi F, Wang ES, et al. Results from a randomized trial of salvage chemotherapy followed by lestaurtinib for patients with FLT3 mutant AML in first relapse. *Blood*. 2011;117(12):3294–3301.
- 36. Röllig C, Serve H, Huttmann A, et al.; Study Alliance Leukaemia. Addition of sorafenib versus placebo to standard therapy in patients aged 60 years or younger with newly diagnosed acute myeloid leukaemia (SORAML): a multicentre, phase 2, randomised controlled trial. *Lancet Oncol.* 2015;16(16):1691–1699.
- 37. Stone RM, Mandrekar S, Sanford BL, et al. The multi-kinase inhibitor midostaurin (M) prolongs survival compared with placebo (P) in combination with daunorubicin (D)/cytarabine (C) induction (ind), high-dose C consolidation (consol), and as maintenance (maint) therapy in newly diagnosed acute myeloid leukemia (AML) patients (pts) age 18-60 with *FLT3* mutations (muts): an international prospective randomized (rand) P-controlled double-blind trial (CALGB 10603/RATIFY [alliance]). *Blood.* 2015;126(23):6.
- 38. Creutzig U, Kaspers GJ. Revised recommendations of the International Working Group for diagnosis, standardization of response criteria, treatment outcomes, and reporting standards for therapeutic trials in acute myeloid leukemia. *J Clin Oncol.* 2004;22(16):3432–3433.
- Stone RM, Mandrekar SJ, Sanford BL, et al. Midostaurin plus chemotherapy for acute myeloid leukemia with a FLT3 mutation. *N Engl J Med.* 2017;377(5):454–464.
- Smith CC, Shah NP. The role of kinase inhibitors in the treatment of patients with acute myeloid leukemia. *Am Soc Clin Oncol Educ Book*. 2013;2013:313–318.
- Röllig C, Bornhauser M, Thiede C, et al. Long-term prognosis of acute myeloid leukemia according to the new genetic risk classification of the European LeukemiaNet recommendations: evaluation of the proposed reporting system. J Clin Oncol. 2011;29(20):2758–2765.
- Dohner H, Estey E, Grimwade D, et al. Diagnosis and management of AML in adults: 2017 ELN recommendations from an international expert panel. *Blood*. 2017;129(4):424–447.

- Best-Aguilera C, Rodrigo GO, Elizabeth GA, Monserrat RR. Treatment of acute myeloid leukemia with the FLT3 gene mutation. *Curr Oncol Rep.* 2017;19(3):21.
- 44. Wander SA, Levis MJ, Fathi AT. The evolving role of FLT3 inhibitors in acute myeloid leukemia: quizartinib and beyond. *Ther Adv Hematol*. 2014;5(3):65–77.
- 45. Stone RM, DeAngelo DJ, Klimek V, et al. Patients with acute myeloid leukemia and an activating mutation in FLT3 respond to a small-molecule FLT3 tyrosine kinase inhibitor, PKC412. *Blood*. 2005;105(1):54–60.
- 46. Döhner H, Perl A, Rousselot P, et al. Efficacy and safety of quizartinib (AC220) in patients age ≥60 years with FLT3-ITD-positive relapsed/ refractory acute myeloid leukemia (AML). Haematologica. 2013;98:233.
- 47. Levis MJ, Martinelli G, Perl AE, et al. The benefit of treatment with quizartinib and subsequent bridging to HSCT for FLT3-ITD(+) patients with AML. *J Clin Oncol*. 2014;32(15_suppl):7093.
- Cortes JE, Kantarjian HM, Kadia TM, et al. Crenolanib besylate, a type I pan-FLT3 inhibitor, to demonstrate clinical activity in multiply relapsed FLT3-ITD and D835 AML. J Clin Oncol. 2016;34(15_suppl):7008.
- 49. Randhawa JK, Kantarjian HM, Borthakur G, et al. Results of a phase II study of crenolanib in relapsed/refractory acute myeloid leukemia patients (Pts) with activating FLT3 mutations. *Blood*. 2014;124(21):389.
- Ohanian M, Kantarjian HM, Borthakur G, et al. Efficacy of a type I FLT3 inhibitor, crenolanib, with idarubicin and high-dose ara-C in multiply relapsed/refractory FLT3+AML. *Blood*. 2016;128(22):2744.
- 51. Levis M, Cortes J, Perl A, et al. High response rate and bridging to hematopoietic stem cell transplantation with quizartinib (AC220) in patients with FLT3-ITD-positive relapsed/refractory acute myeloid leukemia (AML). *Haematologica*. 2013;98:17.
- Faderl S, Wetzler M, Rizzieri D, et al. Clofarabine plus cytarabine compared with cytarabine alone in older patients with relapsed or refractory acute myelogenous leukemia: results from the CLASSIC I trial. *J Clin Oncol.* 2012;30(20):2492–2499.
- Ramos NR, Mo CC, Karp JE, Hourigan CS. Current approaches in the treatment of relapsed and refractory acute myeloid leukemia. *J Clin Med.* 2015;4(4):665–695.
- Sato T, Yang X, Knapper S, et al. FLT3 ligand impedes the efficacy of FLT3 inhibitors in vitro and in vivo. Blood. 2011;117(12):3286–3293.
- Perl AE, Dohner H, Rousselot PH, et al. Efficacy and safety of quizartinib (AC220) in patients age 70 years with FLT3-ITD positive or negative relapsed/refractory acute myeloid leukemia (AML). *J Clin Oncol.* 2013;31(15_suppl):7023.

Supplementary materials

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias	
Knapper et al (2017) ¹	+	•	•	•	+	+	•	
Levis et al (2011) ²	?	+	•	+	+	+	•	
Röllig et al (2015) ²⁶	+	+	•	•	•	+	•	
Serve et al (2013) ⁴	+	•	+	•	•	+	•	
Stone et al (2017) ⁵	+	•	+	+	+	+	+	



Figure SI Risk of bias graph for five RCTs in Part I. Abbreviation: RCT, randomized controlled trial.



Figure S2 Funnel plot for CR rate (A) and ORR (B) compared between FLT3–ITD positive and negative patients. Abbreviations: CR, complete remission; ORR, overall response rate; FLT3, FMS-like tyrosine kinase 3; ITD, internal tandem duplication; SE, standard error; OR, odds ratio.

Study	Selection				Comparability	Outcome	Score		
(reference)	Representa- tiveness of exposed cohort	Selection of nonexposed cohort	Ascertainment of exposure	Outcome of interest not present at start		Assessment of outcome	Follow- up length	Follow- up adequacy	
Alvarado et al ⁶	*	*	*		*	*	*		6
Cooper et al ⁷		*	*		**	*	*		6
Cortes et al ⁸		*	*		*	*	*	*	6
Cortes et al ⁹		*	*		**	*	*	*	7
Crump et al ¹⁰	*	*	*	*	*	*	*		7
Fiedler et al ¹¹		*	*		**	*	*	*	7
Fiedler et al ¹²		*	*		**	*	*	*	7
Fischer et al ¹³ *		*	*		**	*	*	*	7
Inaba et al ¹⁴		*	*		**	*	*	*	7
Knapper et al ²⁴		*	*	*	**	*	*	*	8
Levis et al ²³		*	*		*	*	*	*	6
Macdonald et al ¹⁵		*	*		**	*	*	*	7
Pratz et al ¹⁶		*	*		*	*	*	*	6
Ramsingh et al ¹⁷		*	*		**	*	*		6
Ravandi et al ¹⁸		*	*	*	**	*	*	*	8
Ravandi et al ¹⁹		*	*	*	**	*	*	*	8
Stone et al ²⁵		*	*	*	**	*	*	*	8
Walker et al ²⁰		*	*	*	**	*	*		7
Röllig et al ³	*	*	*		**	*	*		7
lyer et al ²¹		*	*		*	*	*	*	6
Zhang et al ²²		*	*		*		*	*	5

Table SI Assessment of methodological quality of cohort studies

Note: NOS: study can have one point (*) for meeting each criterion, except that comparability can have a maximum of two points (**). Abbreviation: NOS, Newcastle-Ottawa Quality Assessment Scale.

References

- Knapper S, Russell N, Gilkes A, et al. A randomized assessment of adding the kinase inhibitor lestaurtinib to first-line chemotherapy for FLT3-mutated AML. *Blood.* 2017;129(9):1143–1154.
- 2. Levis M, Ravandi F, Wang ES, et al. Results from a randomized trial of salvage chemotherapy followed by lestaurtinib for patients with FLT3 mutant AML in first relapse. *Blood*. 2011;117(12):3294–3301.
- Rollig C, Brandts C, Shaid S, et al. Survey and analysis of the efficacy and prescription pattern of sorafenib in patients with acute myeloid leukemia. *Leuk Lymphoma*. 2012;53(6):1062–1067.
- Serve H, Krug U, Wagner R, et al. Sorafenib in combination with intensive chemotherapy in elderly patients with acute myeloid leukemia: results from a randomized, placebo-controlled trial. *J Clin Oncol.* 2013;31(25):3110–3118.
- Stone RM, Mandrekar SJ, Sanford BL, et al. Midostaurin plus chemotherapy for acute myeloid leukemia with a FLT3 mutation. N Engl J Med. 2017;377(5):454–464.
- Alvarado Y, Kantarjian HM, Luthra R, et al. Treatment with FLT3 inhibitor in patients with FLT3-mutated acute myeloid leukemia is associated with development of secondary FLT3-tyrosine kinase domain mutations. *Cancer*. 2014;120(14):2142–2149.
- Cooper TM, Cassar J, Eckroth E, et al. A phase I study of quizartinib combined with chemotherapy in relapsed childhood leukemia: a therapeutic advances in childhood leukemia & lymphoma (TACL) study. *Clin Cancer Res.* 2016;22(16):4014–4022.
- Cortes JE, Perl AE, Dombret H, et al. Final results of a phase 2 open-label, monotherapy efficacy and safety study of quizartinib (AC220) in patients 60 years of age with FLT3 ITD positive or negative relapsed/refractory acute myeloid leukemia. *Blood.* 2012;120 (21):48.

- 9. Cortes JE, Kantarjian H, Foran JM, et al. Phase I study of quizartinib administered daily to patients with relapsed or refractory acute myeloid leukemia irrespective of FMS-like tyrosine kinase 3-internal tandem duplication status. *J Clin Oncol.* 2013;31(29):3681–3687.
- Crump M, Hedley D, Kamel-Reid S, et al. A randomized phase I clinical and biologic study of two schedules of sorafenib in patients with myelodysplastic syndrome or acute myeloid leukemia: a NCIC (National Cancer Institute of Canada) Clinical Trials Group Study. *Leuk Lymphoma*. 2010;51(2):252–260.
- Fiedler W, Serve H, Dohner H, et al. A phase 1 study of SU11248 in the treatment of patients with refractory or resistant acute myeloid leukemia (AML) or not amenable to conventional therapy for the disease. *Blood*. 2005;105(3):986–993.
- Fiedler W, Kayser S, Kebenko M, et al. A phase I/II study of sunitinib and intensive chemotherapy in patients over 60 years of age with acute myeloid leukaemia and activating FLT3 mutations. *Br J Haematol.* 2015;169(5):694–700.
- 13. Fischer T, Stone RM, Deangelo DJ, et al. Phase IIB trial of oral midostaurin (PKC412), the FMS-like tyrosine kinase 3 receptor (FLT3) and multi-targeted kinase inhibitor, in patients with acute myeloid leukemia and high-risk myelodysplastic syndrome with either wild-type or mutated FLT3. J Clin Oncol. 2010;28(28):4339–4345.
- Inaba H, Rubnitz JE, Coustan-Smith E, et al. Phase I pharmacokinetic and pharmacodynamic study of the multikinase inhibitor sorafenib in combination with clofarabine and cytarabine in pediatric relapsed/ refractory leukemia. *J Clin Oncol.* 2011;29(24):3293–3300.
- Macdonald DA, Assouline SE, Brandwein J, et al. A phase I/II study of sorafenib in combination with low dose cytarabine in elderly patients with acute myeloid leukemia or high-risk myelodysplastic syndrome from the National Cancer Institute of Canada Clinical Trials Group: trial IND186. *Leuk Lymphoma*. 2013;54(4):760–766.

- Pratz KW, Cho E, Levis MJ, et al. A pharmacodynamic study of sorafenib in patients with relapsed and refractory acute leukemias. *Leukemia*. 2010;24(8):1437–1444.
- Ramsingh G, Westervelt P, McBride A, et al. Phase I study of cladribine, cytarabine, granulocyte colony stimulating factor (CLAG regimen) and midostaurin and all-trans retinoic acid in relapsed/refractory AML. *Int J Hematol.* 2014;99(3):272–278.
- Ravandi F, Cortes JE, Jones D, et al. Phase I/II study of combination therapy with sorafenib, idarubicin, and cytarabine in younger patients with acute myeloid leukemia. *J Clin Oncol.* 2010;28(11):1856–1862.
- Ravandi F, Arana YC, Cortes JE, et al. Final report of phase II study of sorafenib, cytarabine and idarubicin for initial therapy in younger patients with acute myeloid leukemia. *Leukemia*. 2014;28(7):1543–1545.
- Walker AR, Wang H, Walsh K, et al. Midostaurin, bortezomib and MEC in relapsed/refractory acute myeloid leukemia. *Leuk Lymphoma*. 2016;57(9):2100–2108.
- Iyer SP, Jethava Y, Karanes C, Eckardt JR, Collins R. Safety study of salvage chemotherapy high-dose ara-C/mitoxantrone (HAM) and type I FLT3-TKI crenolanib in first relapsed/primary refractory AML. *Blood*. 2016;128(22):3983–3983.

- Zhang W, Konopleva M, Shi YX, et al. Mutant FLT3: a direct target of sorafenib in acute myelogenous leukemia. *J Natl Cancer Inst.* 2008;100(3):184–198.
- 23. Levis MJ, Perl AE, Dombret H, et al. Final results of a phase 2 openlabel, monotherapy efficacy and safety study of quizartinib (AC220) in patients with FLT3-ITD positive or negative relapsed/refractory acute myeloid leukemia after second-line chemotherapy or hematopoietic stem cell transplantation. *Blood*. 2012;120(21):673.
- 24. Knapper S, Burnett AK, Littlewood T, et al. A phase 2 trial of the FLT3 inhibitor lestaurtinib (CEP701) as first-line treatment for older patients with acute myeloid leukemia not considered fit for intensive chemotherapy. *Blood.* 2006;108(10):3262–3270.
- 25. Stone RM, Fischer T, Paquette R, et al. Phase IB study of the FLT3 kinase inhibitor midostaurin with chemotherapy in younger newly diagnosed adult patients with acute myeloid leukemia. *Leukemia*. 2012;26(9):2061–2068.
- 26. Röllig C, Serve H, Huttmann A, et al.; Study Alliance Leukaemia. Addition of sorafenib versus placebo to standard therapy in patients aged 60 years or younger with newly diagnosed acute myeloid leukaemia(SORAML): a multicentre, phase 2, randomised controlled trial. *Lancet Oncol.* 2015;16(16):1691–1699.

Cancer Management and Research

Publish your work in this journal

Cancer Management and Research is an international, peer-reviewed open access journal focusing on cancer research and the optimal use of preventative and integrated treatment interventions to achieve improved outcomes, enhanced survival and quality of life for the cancer patient. The manuscript management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit http://www.dovepress.com/testimonials.php to read real quotes from published authors.

Submit your manuscript here: https://www.dovepress.com/cancer-management-and-research-journal

Dovepress