

Review Article

Prognostic Value of Circulating Tumor DNA (ctDNA) in Diffuse Large B-Cell Lymphoma (DLBCL): Systemic Review and Meta-Analysis.

Anu Kumari¹, PhD, Dr. Anupriya Kaur², Charanpreet Singh³, Dr. Anju Goyal⁴, Dr. Priyanka Srivastava⁵, Dr. Arihant Jain⁶, Dr. Sreejesh Sreedharanunni⁷, Dr. Amanjit Bal⁸, Dr. Alka Rani Khadwal⁹, Dr. Pankaj Malhotra¹⁰.

1. Department of Clinical Hematology & Medical oncology, PGIMER, Chandigarh. **Email:** anu1995gn@gmail.com
2. (Associate Professor), Genetics & Metabolic Unit, Advanced Pediatric Centre, PGIMER, Chandigarh. **Email:** anukaur.genetics@gmail.com
3. Department of Clinical Hematology & Medical oncology, PGIMER, Chandigarh. **Email:** charanpreetsingh14@gmail.com
4. Assistant Professor, Department of statistics, Panjab University, Chandigarh. **Email:** anju.statistics@gmail.com
5. Assistant Professor, Genetics & Metabolic Unit, Advanced Pediatric Centre, PGIMER, Chandigarh. **Email:** srivastavapriy@gmail.com
6. (Associate Professor), Department of Clinical Hematology & Medical Oncology, PGIMER, Chandigarh, **Email:** drarihantjain86@gmail.com
7. Additional Professor, Department of Hematology, PGIMER, Chandigarh, **Email:** dr.s.sreejesh@gmail.com
8. Professor, Department of Histopathology, PGIMER, Chandigarh, **Email:** docaman6@gmail.com
9. (Professor), Department of Clinical Hematology & Medical Oncology, PGIMER, Chandigarh, **Email:** alkakhadwal@hotmail.com
10. Professor & HOD, Department of Clinical Hematology & Medical Oncology, PGIMER, Chandigarh, **Email:** malhotrapankaj@hotmail.com

Abstract

Circulating cell-free DNA (ccf-DNA) and circulating tumor DNA (ctDNA) provide a minimally invasive method for cancer detection and measurement. However, their diagnostic and prognostic significance in hematological malignancies remains ambiguous. This meta-analysis aims to evaluate the prognostic value of ccf-DNA or ctDNA in patients with diffuse large B-cell lymphoma (DLBCL). All relevant literature was retrieved through a systematic search of electronic databases, including PubMed, Embase, Scopus, and the Cochrane Library. Eight eligible studies were selected for the analysis of prognostic value of ccf-DNA or ctDNA. Statistical analyses were performed using R software. The results indicate significant associations with both PFS (HR = 2.14; 95% CI: 1.31–3.40; $p < 0.01$) and OS (HR = 2.51; 95% CI: 1.84–3.40) for patients with elevated ccf-DNA or ctDNA levels. The results of this meta-analysis strongly suggest that elevated levels of ccfDNA or ctDNA are indicative of poor prognosis in patients with DLBCL.

Keywords : Circulating cell-free DNA, circulating tumor DNA, DLBCL, Prognosis, Hazard Ratio, Meta-analysis.

INTRODUCTION

Diffuse large B-cell lymphoma (DLBCL) is the most common subtype of lymphoma, currently, the International Prognostic Index (IPI) is used for risk stratification of DLBCL which is an old prognostic score with limitations and there is an unmet need for a more refined prognostic tool with better representation of tumor biology (1). Over the past decade, liquid biopsy techniques utilizing circulating cell-free DNA (ccf-DNA) or circulating tumor DNA (ctDNA) from peripheral blood have emerged as robust diagnostic and prognostic tools for various cancers, including lung, prostate, and

gastrointestinal cancers (2-4).

The role of ccf-DNA or ctDNA as a tool for risk stratification in patients with aggressive lymphoma has gained attention in recent years. However, most studies in this area are small, single-center experiences, and their findings are varied (5-12). There is limited systematic evidence demonstrating the prognostic value of ccf-DNA or ctDNA for DLBCL patients treated with R-CHOP regimen (13-15).

To address this gap, we conducted a systematic review and meta-analysis of studies that utilized ccf-DNA or ctDNA for prognostication of DLBCL, either at the time of diagnosis or during the course of treatment.

***Corresponding Author:** Dr. Gaurav Prakash, Professor, Department of Clinical Hematology and Medical Oncology, Post-Graduate Institute of Medical Education and Research, India, **E-mail:** drgp04@gmail.com.

Received: 20-April-2025, Manuscript No. TEJOC- 4773 ; **Editor Assigned:** 22-April-2025 ; **Reviewed:** 14-May-2025, QC No. TEJOC- 4773 ;

Published: 06-June-2025, **DOI:** 10.52338/tejoc.2025.4773

Citation: Dr. Gaurav Prakash. Prognostic value of circulating tumor DNA (ctDNA) in Diffuse Large B-Cell Lymphoma (DLBCL): Systemic review and meta-analysis. The European Journal of Cancer. 2025 June; 11(1). doi: 10.52338/tejoc.2025.4773.

Copyright © 2025 Dr. Gaurav Prakash. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

MATERIALS AND METHODS

This meta-analysis was conducted according to the guidelines provided by the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA). The study was also registered with the International Prospective Register of Systematic Reviews (PROSPERO) with registration number CRD42024520358.

Inclusion criteria

1. Patients with treatment naïve DLBCL
2. Studies reporting ccf-DNA or ctDNA in plasma/serum at one or more than one time points
3. Studies with information on patients' survival outcomes such as OS, PFS and EFS
4. Studies mentioning the technique for ccf-DNA or ctDNA detection or quantification.

Exclusion criteria

1. All case reports, case series, review articles, editorials, letters or comments, and conference abstracts were excluded
2. Studies without survival data/Hazard ratio (HR) required to perform meta-analysis were excluded
3. Duplicate studies/ abstracts were excluded
4. Studies published in non-English language were excluded

Search Strategy

Literature searches were conducted by searching electronic databases of Pubmed, Embase, Scopus and Cochrane Library for relevant papers published up to 31 May 2024. The following terms were used for searching: "ctDNA" or "cell-free DNA" or "cell-free tumor DNA" or "Circulating Nucleic Acids" or "Cell-Free Deoxyribonucleic acid" or "Cell-Free Nucleic Acid" or "circulating tumour DNA" or "tumor DNA" and "non-Hodgkin lymphoma" or "NHL" or "lymphoma". The Mesh terms and detailed search strategy is provided in the supplement file.

Study Screening and Selection

Relevant articles were selected based on predefined eligibility criteria, and studies not meeting these criteria were excluded from the analysis. Two independent reviewers thoroughly analyzed and assessed the texts of the selected articles to confirm their eligibility before extracting data. Additionally, a manual search of the reference lists of the selected articles was conducted to ensure the inclusion of any relevant studies that were not initially identified through the search

Data extraction and quality assessment

Two independent reviewers performed data extraction. The following variables were extracted from each study: first author, year of publication, country, study design, number of

participants, age, sex, disease stage, bulky/non-bulky disease, IPI, PS, LDH, ccf-DNA or ctDNA measurement method, target gene(s), time point(s) of ccf-DNA or ctDNA measurement, follow up period of the study and HR value of prognostic index with 95% confidence intervals (CI) for survival outcomes. Extracted data was entered in an excel sheet and analyzed for outcomes. The Newcastle-Ottawa Scale (NOS) checklist was used to evaluate the quality of the eligible studies.

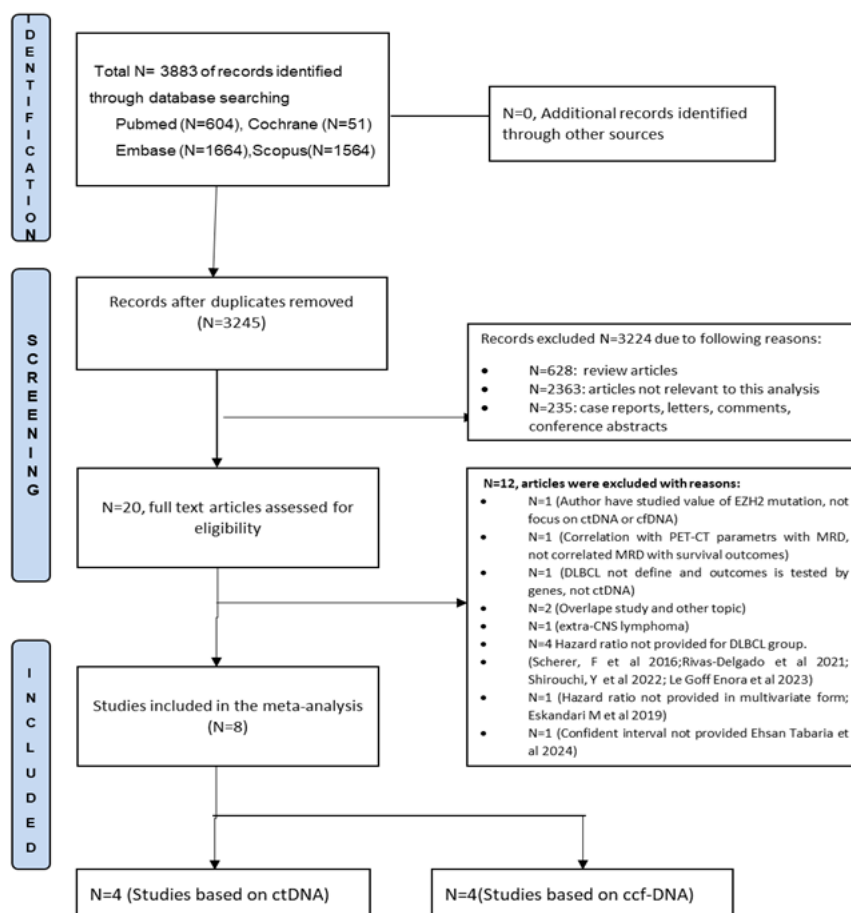
Data synthesis and Statistical analysis

The total HR was computed with the addition of each study. Heterogeneity among the studies was assessed using the I^2 statistic. An I^2 value exceeding 40% indicated significant heterogeneity. In cases of significant heterogeneity, a random-effects (RE) model was employed; otherwise, a fixed-effects (FE) model was used. A pooled HR of ≥ 1 indicated a worse survival outcome for patients with detectable ccf-DNA or ctDNA. Subgroup analyses were conducted to explore sources of heterogeneity between studies. Publication bias was assessed by examining the asymmetry of the funnel plot, and the Egger test was used to detect publication bias among the studies. Statistical analyses were performed using R software, with a P-value of < 0.05 considered statistically significant.

RESULTS

Literature screening and selection of study

In the process of literature screening and study selection, 3883 potentially relevant articles were identified and after removing duplicates, 3245 records were selected for further study. Three thousand, two hundred, twenty four records were excluded because they were reviews, case reports, letters to the editor, non-English articles, or conference abstracts. This screening process ensured that only relevant studies, according to the inclusion criteria, were included for subsequent analysis. Full text of 20 studies was reviewed, and 12 studies were excluded due to reasons of lack of focus on ccf-DNA or ctDNA or absence of HR for outcome. Finally, 8 eligible studies were selected for meta-analysis, encompassing 427 patients with baseline ccf-DNA or ctDNA values and 287 patients with ccf-DNA or ctDNA values measured after starting anticancer therapy. The PRISMA flow chart presenting the steps of the study selection in detail is shown in **Figure (1)**.

Figure 1. PRISMA flow diagram of the study selection process.

Data extraction and quality management

Data extraction and quality assessment were conducted across all selected studies, which were published between 2016 and 2024. Each of the eight studies evaluated ccf-DNA or ctDNA in plasma (5-6,8-12) with one study assessing ctDNA both in serum and plasma (7) (**Table 1**). Out of eight selected studies, four studies focused on ctDNA and another four studies focused on ccf-DNA values (**Table 2**). Two studies provided ccf-DNA values, measured in ng/ml, one study utilized methylation levels for ccf-DNA quantification, and one study used a 29-gene-based weighted prognostic score (wp-score) to measure ctDNA levels. Four studies used a threshold based on log haploid genome equivalent per ml (hGE/mL) or Variant Allele Frequency (VAF) (**Table 2**).

The hazard ratio for outcomes, accompanied by their 95% confidence intervals (95% CI), were directly extracted from selected studies. Five studies examined prognostic value of ccf-DNA or ctDNA on the basis of baseline values (5-9) while, two studies utilized ctDNA values after starting anticancer therapy (but before completion of treatment) (10, 12), one study has estimate ctDNA both at baseline and after initiation of therapy (11) (**Table 2**).

On the Newcastle-Ottawa Scale (NOS), notably, all studies achieved scores exceeding seven, indicating robust methodological quality and reliant ability. Detailed are available in **Table 3**.

Table 1. Characteristics of included studies for quantitative synthesis of meta-analysis.

S.No.	Author	Country	Type of study	Patients	Patients number	Type of treatment	Source of sample	Time point of plasma / serum collection	Technique	Gene Symbol	Follow-up time period
1	Kristensen, L S et al., 2016	Denmark	Retro	DLBCL	71	59 received RCHOP-like chemotherapy FOR others not reported	Plasma	Baseline	PCR: Pyrosequencing	DAPK1	NR
2	Li M et al., 2017	China	Pros	DLBCL	98	Chemotherapy type not mentioned	Plasma	Baseline	Fluorometer, qPCR	APP Gene	13.5 (1 - 36 months)
3	Kurtz, D. M et al., 2018	North America and Europe	Retro	DLBCL	108	RCHOP-45%; EPOCH R 34%;Others-22%	Plasma/ Serum	Baseline	CAPP-Seq NGS	TP53,BCL2, BCL6,CARD11 etc.	31.2 months
4	Chiu, B.C.-H. et al., 2018	America	Pros	DLBCL	48	RCHOP- 66.6%; EPOCH R- 16.7%	Plasma	Baseline	Illumina NextSeq 500 platform (NGS)	5hmC	NR
5	Hur, J Y et al.,2020	Korea	Pros	DLBCL	51	Chemotherapy type not mentioned	Plasma	Baseline	Qubit 2.0 Fluorometer;2200 TapeStation Instrument	NA	NR
6a	Guan, T et al., 2022	China	Pros	DLBCL- Training	85	RCHOP	Plasma	within 1week of receiving anticancer treatment and	NGS: Illumina NovaSeq 5000	59 gene Panel	NR
6b	Guan, T et al., 2022	China	Pros	DLBCL- Validation	84	RCHOP	Plasma	within 1week of receiving anticancer treatment and	NGS: Illumina NovaSeq 5000	60 gene Panel	NR
7a	Li, M et al., 2022	China	Pros	DLBCL	51	R-CHOP or R-CHOP like	Plasma	Baseline	NovaSeq6000 NGS platform (Illumina)	188 Gene Panel	30.3 (range, 3.8–101.2) months
7b	Li, M et al., 2022	China	Pros	DLBCL	37	R-CHOP or R-CHOP like	Plasma	After 2 cycles of therapy	NovaSeq6000 NGS platform (Illumina)		
7c	Li, M et al., 2022	China	Pros	DLBCL	38	R-CHOP or R-CHOP like	Plasma	After 4 cycles of therapy	NovaSeq6000 NGS platform (Illumina)		
8	Miguel Alcoceba et al., 2022	Spain	Pros	DLBCL	43	R-CHOP or R-CHOP like	Plasma	After 2 cycles of therapy	NGS:NextSeq 500 (Illumina)	Panel	3 year (range 0.2–4.8)

Table 2. Survival Outcomes of included studies for quantitative synthesis of the meta-analysis.

S.No.	Author	Time point of plasma /serum collection	Survival Outcome	Defination of cfDNA positive	ctDNA +ve	PFS/EFS(HR)	OS (HR)
1	Kristensen, L S et al., 2016	Baseline	OS	Aberrant methylation level; The cutoffs were 5.5% for DAPK1.	14 (19%)		8.90 (2.70–29.30), p=0.0007
2	Li M et al., 2017	Baseline	PFS	1586 ng/ml cfDNA	NA	1.45 (0.490–4.263), p=0.504	
3	Kurtz, D. M et al., 2018	Baseline	OS;EFS	>2.5 log hGE/mL threshold	212 (98%)	1.90 (1.12-3.23)	1.30 (0.65-2.59), p=0.46
4	Chiu, B.C.-H. et al., 2018	Baseline	EFS	29 gene based weighted prognostic score (wp-score) High/Low	23 High wp score , 23 Low wp score	9.17 (2.01–41.89), p=0.04	
5	Hur, J Y et al.,2020	Baseline	OS;EFS	> 16.8 ng/ml cfDNA for EFS, and >30.3 ng/ml cfDNA for OS	19 High cfDNA; 32 Low cfDNA	5.33 (1.72–16.52), p=0.003	4.51 (1.35-15.05), p=0.014
6a	Guan, T et al., 2022	within 1 week of receiving anticancer treatment and	PFS;OS	a mean VAF value $\geq$ 4.94%,	ctDNA positive in 64 (75.3%) patients	2.5 (1.1-5.8), p=0.04	2.6 (1.1-6.4), p=0.04
6b	Guan, T et al., 2022	within 1 week of receiving anticancer treatment and	PFS;OS	a mean VAF value $\geq$ 4.94%,	ctDNA positive in 67 (79.8%) patients	1.8 (0.8-4.0), p=0.014	1.9 (0.8-4.3), p=0.093
7a	Li, M et al., 2022	Baseline	PFS;OS	.>2.44 log hGE/ ml(Concentrations of ctDNA were expressed in log hGE/ mL,	High=26 ; Low=25	2.47 (1.35-4.5), p=0.004	2.49 (1.238-5), p=0.011
7b	Li, M et al., 2022	After 2 cycles of therapy	PFS;OS	.>2.44 log hGE/ ml(Concentrations of ctDNA were expressed in log hGE/ mL,	High=21 ; Low=16	2.22 (1.14-4.3), p=0.019	2.79 (1.20-6.5), p=0.017
7c	Li, M et al., 2022	After 4 cycles of therapy	PFS;OS	ctDNA negative or positive	28=ctDNA negative 10= ctDNA postive	3.65 (1.452-9.178, p=0.0003	3.56 (1.192-10.49), p=0.016
8	Miguel Alcoceba et al., 2022	After 2 cycles of therapy	PFS	2.5 log hGE/mL	NA	0.193 [0.058–0.639], p=0.007	

Table3. Quality assessment of all included studies.

S.No.	Study ID	Selection (Maximum 4 points)				Comparability (Maximum 2 points)	Outcome (Maximum 3 points)			Total points (Maximum 9 points)
		Representative of exposed cohort	Selection of the non- exposed cohort	Ascertainment of exposure	Outcomes not presented at start of study	Impact of bias	Assessment of outcome	Was follow- up long enough for outcomes occur	Adequacy of follow-up	
1	Kristensen, L S et al., 2016	*	*	*	*	*	*	*	*	8
2	Li M et al., 2017	*	*	*	*	*	*	*	*	8
3	Kurtz, D. M et al., 2018	*	*	*	*	*	*	*	*	8
4	Chiu, B.C.H. et al., 2019	*	*	*	*	*	*	*	*	8
5	Hur, J Y et al. 2020	*	*	*	*	*	*	*	*	8
6	Guan, T et al., 2022	*	*	*	*	*	*	*	*	8
7	Li, M et al. 2022	*	*	*	*	*	*	*	*	8
8	Miguel Alcoceba et al., 2024	*	*	*	*	*	*	*	*	8

Techniques used for Detection of ctDNA or ccf-DNA

Detection of ccf-DNA or ctDNA in the reviewed studies predominantly utilized Polymerase Chain Reaction (PCR) and Next-Generation Sequencing (NGS) techniques. Among the 8 studies included, NGS was employed in 5 studies (7-8, 10-12), PCR in one study (5) and fluorometry in two studies (6, 9) . Specifically, out of the 5 studies using NGS technology, one study employed Cancer Personalized Profiling by Deep Sequencing (CAPP-Seq) to measure ctDNA levels (7) , while the other 4 studies utilized the Illumina platform for sequencing (8,10-12) . The 8 studies investigated PFS, EFS, and OS as outcome indicators using multivariate analysis. These data were pooled for meta-analysis to examine the association between ccf-DNA or ctDNA levels and the prognosis of patients with DLBCL. The results of this meta-analysis are illustrated in **Figure 2, 3 and 4** depicting synthesized evidence and statistical outcomes derived from the combined analysis of the selected studies. These figures typically present key metrics such as HR with 95% confidence intervals (CIs) and forest plots to visualize the pooled effect sizes and their variability across studies.

Prognostic role of ccf-DNA or ctDNA

Prognostic value of ccf-DNA or ctDNA of all eight studies was analyzed. The results indicate significant associations with both PFS and OS. Specifically, for patients with high ccf-DNA or ctDNA, HR for PFS was 2.14 (95% CI, 1.31–3.40) (Figure 2C), and for OS 2.51 (95% CI, 1.84–3.40) (Figure 3C). Significant heterogeneity was observed in data analysis for PFS ($I^2 = 62\%$, $p < 0.01$) (Figure 2C), indicating variability across the studies included in the meta-analysis. Therefore, a random-effects model was employed to account for this heterogeneity and provide a more conservative estimation of the pooled effect size. For OS, no significant heterogeneity was observed ($I^2 = 27\%$, $p = 0.21$) thus a fixed effect model was applied (Figure 3C).

Figure 2. Forest plot of HR for ccf-DNA/ctDNA associated with PFS in DLBCL: A) the original HRs of PFS for ccf-DNA/ctDNA before initiation of therapy. B) The original HRs of PFS for ccf-DNA/ctDNA after starting anticancer therapy. C) The original HRs of PFS for ccf-DNA/ctDNA for all studies.

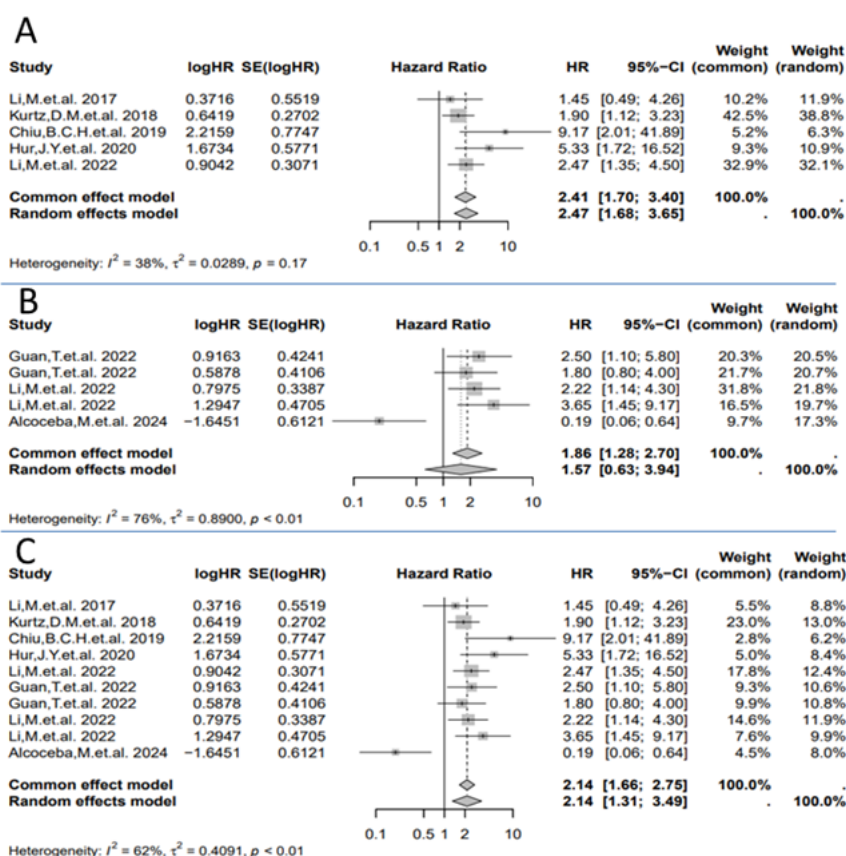


Figure 3. Forest plot of HR for ccf-DNA/ctDNA associated with OS in DLBCL: A) the original HRs of OS for ccf-DNA/ctDNA at baseline. B) The original HRs of OS for ccf-DNA/ctDNA after starting anticancer therapy. C) The original HRs of OS for ccf-DNA/ctDNA for all studies.

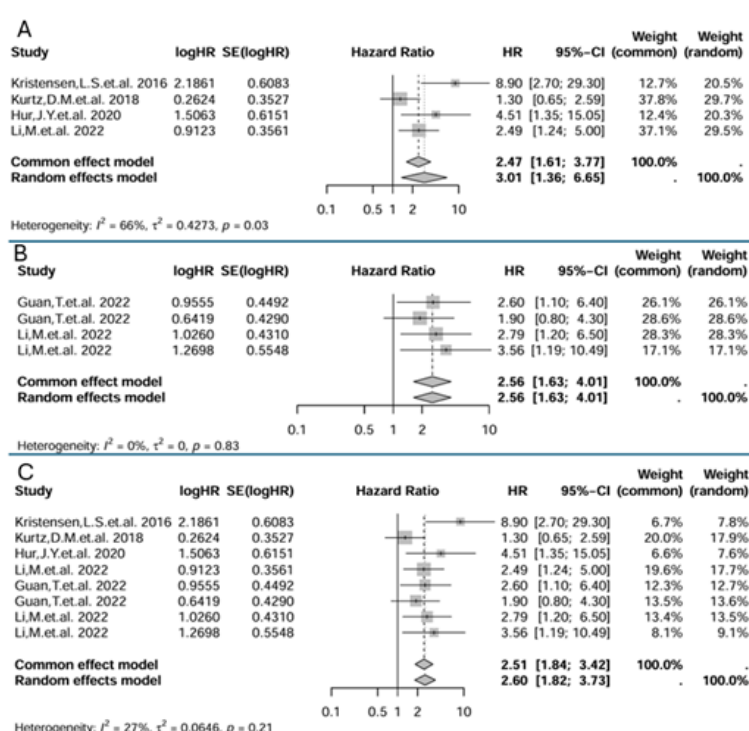
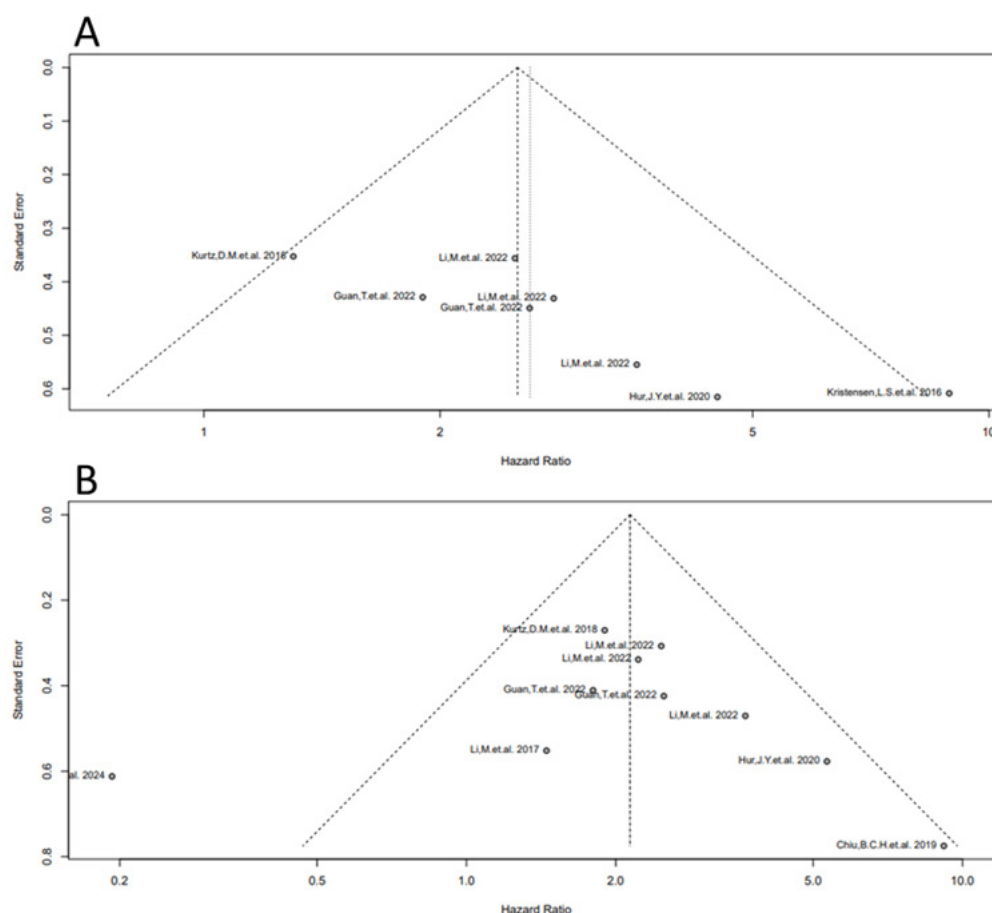


Figure 4. Funnel plot for the assessment of publication bias of the included studies. A) Funnel plot of OS for all included studies B) Funnel plot of PFS for all included studies.



Subgroup analysis

Subgroup analysis was performed to study difference in prognostic value of ccf-DNA or ctDNA according to patients' ethnicity, method of ccf-DNA or ctDNA estimation, time of ccf-DNA or ctDNA sampling with respect to administration of chemotherapy, and type of DNA studied (ccf-DNA vs ctDNA) (supplement file).

Prognostic role of ccf-DNA or ctDNA at baseline

Five studies have assessed baseline levels of ccf-DNA or ctDNA, revealing that patients with elevated levels prior to therapy had significantly worse OS (HR = 3.01; 95% CI, 1.36–6.65, $p = 0.03$) and PFS (HR = 2.41; 95% CI, 1.70–3.40, $p = 0.17$).

Prognostic role of only ccf-DNA at Baseline

In three studies that utilized PFS as the outcome measure, elevated levels of ccf-DNA were associated with poor PFS (HR = 3.87; 95% CI: 1.30–11.48). Furthermore, in two studies, high levels of ccf-DNA were correlated with poor OS (HR = 6.36; 95% CI: 2.72–14.85). No studies were found which measured ccf-DNA during or after therapy.

To further analyze the relationship between ctDNA and survival in patients with DLBCL, we conducted additional subgroup analyses focusing exclusively on studies that evaluated ctDNA.

Prognostic role of ctDNA at Baseline

In two studies, elevated levels of ctDNA were associated with poor PFS (HR = 2.13; 95% CI: 1.43–3.17) and poor OS (HR = 1.80; 95% CI: 0.95–3.40).

Prognostic role of ctDNA at ongoing therapy and after therapy

Several studies assessed ctDNA following therapy, revealing that high ctDNA levels were linked to poor PFS according to three studies (HR = 1.57; 95% CI: 0.63–3.94; $I^2 = 76\%$, $p < 0.01$) and poor OS according to two studies (HR = 2.56; 95% CI: 1.63–4.01; $I^2 = 0\%$, $p = 0.83$).

Publication bias

To assess publication bias, funnel plots were examined for both OS and PFS, revealing a roughly symmetrical shape, as illustrated in Figures 4A and 4B. Additionally, the P value from Egger's test was 0.96. This indicates that there was no significant evidence of publication bias.

DISCUSSION

Most hematologic malignancies benefit from the availability of blood or bone-marrow samples for diagnosis, risk stratification, and monitoring treatment response (16).

However, due to the organ- or tissue-specific presentation of most lymphomas, a blood-based test for diagnosis or monitoring depth-of-response is not currently available. Recently, the use of ccf-DNA or ctDNA as a non-invasive biomarker has gained significant attention for many solid tumors. (17). Developing ccf-DNA or ctDNA as a robust prognostic tool, similar to its use in solid tumors, would be a valuable addition to the management of patients with lymphoma.

We conducted a comprehensive meta-analysis to evaluate the potential significance of ccf-DNA or ctDNA in patients with DLBCL who were treated with combination chemotherapy (mainly R-CHOP or R-CHOP-like therapies) After screening many studies met the inclusion criteria, however they were excluded due to limitations in data availability (18-30) and finally total 8 eligible studies comprising of 714 patients were analyzed (5-12).

The optimal time point for detection of ccf-DNA or ctDNA is not yet defined. We found that five studies analyzed ccf-DNA or ctDNA exclusively at baseline (5-9), two studies only after starting chemotherapy (10,12), and one study assessed ctDNA both at baseline and after initiation of therapy (11). As a result we grouped the studies according to the time of estimation of ccf-DNA or ctDNA.

In five of the selected studies, NGS was used to detect ccf-DNA or ctDNA, (7-8,10-12) whereas in three studies PCR with fluorometric techniques was used for estimation of ccf-DNA (5-6,9). PCR-based methods are both rapid and cost-effective but are limited in their ability to analyze only specific loci at a time. In contrast, NGS-based methods can assess a larger number of loci and even perform whole-exome sequencing, though they generally have lower sensitivity compared to PCR-based methods and are more expensive.

After synthesizing the composite evidence, we found that higher ccf-DNA or ctDNA values before starting treatment were significantly associated with PFS and OS, with HR of 2.41 (1.70-3.40) and 3.01 (1.36-6.65), respectively (31-32). However, testing of ctDNA after initiating antilymphoma therapy did not prognosticate for PFS but was significantly associated with poor OS, with HRs of 1.57 (0.63-3.94) for PFS and 2.56 (1.63-4.01) for OS.

When we combined the data for ccf-DNA or ctDNA assessed at any time point during treatment, higher values were found to be associated with poor PFS and OS, with HRs of 2.14 (1.31-3.49) for PFS and 2.51 (1.84-3.41) for OS (Figure 2,3) (33)

Kurtz et al. (2018) investigated the clinical utility of ctDNA profiling with CAPP-Seq in DLBCL patients, focusing on mutational genotyping and disease burden assessment. Their objective was to evaluate whether ctDNA quantification before and during treatment could predict EFS and OS. The study determined a threshold level of 2.5 log hGE/mL to stratify patients based on EFS. It was found that pretreatment

ctDNA levels had a strong correlation with the IPI and Total Metabolic Tumor Volume (TMTV), indicating that ctDNA could serve as an independent surrogate marker for disease burden beyond these established factors (7).

Miaomiao Li et al. (2022) used a lymphoma-specific sequencing panel and similarly categorized ctDNA levels as log hGE/mL to evaluate the prognostic and predictive value of ctDNA measurements before, during, and after first-line therapy in 73 Chinese DLBCL patients (11) Miguel Alcoceba et al. (2023) also reported ctDNA levels as log hGE/mL, establishing various cutoff points (ranging from 2.0 to 3.5 log hGE/mL) to define the pretreatment ctDNA threshold for predicting PFS (12). These studies collectively highlight the emerging role of ctDNA as a promising biomarker in the management of DLBCL. Incorporating ctDNA analysis into clinical practice could enhance risk stratification, tailor treatment strategies, and ultimately improve patient outcomes in this aggressive form of lymphoma.

Li M et al. (2017) employed PCR and fluorometric technology to evaluate ccf-DNA as a prognostic biomarker and found that high levels of ccfDNA and an elevated integrity index were linked to poor prognosis (6). Similarly, Kristensen et al. (2016) suggested that DAPK1 methylation in ccf-DNA from plasma could serve as a useful biomarker for assessing treatment response in DLBCL (5). Hur et al. (2020) found that high ccf-DNA levels independently predicted poorer EFS in DLBCL patients (9). There is another study conducted by Chiu BC et al., 2019 developed a 29-gene weighted prognostic score (wp-score) using elastic net regularization in a Cox model to predict EFS and OS. Patients with high wp-scores had significantly worse EFS, compared to low-risk patients (8).

The cumulative findings from these studies indicate that ccf-DNA and its methylation status may also substantially improve prognostic accuracy and inform treatment strategies, thereby enhancing patient outcomes. Additionally, these studies underscore the potential of ccf-DNA as a valuable prognostic biomarker in the management of aggressive lymphomas, particularly DLBCL.

Our meta-analysis revealed that patients with higher ccf-DNA or ctDNA at baseline and after the initiation of therapy experienced worse PFS and OS. These results indicate the consistent prognostic value of ctDNA at various time points in DLBCL patients, underscoring its potential as a reliable biomarker for predicting clinical outcomes.

In the three studies that evaluated ccf-DNA at baseline using PFS as the outcome measure, high levels of ccf-DNA were associated with poor PFS (HR = 3.87; 95% CI:1.30-11.48) (6,8-9). Additionally, two studies indicated that elevated ccf-DNA levels correlated with poor OS (HR = 6.36; 95% CI: 2.72-14.85) (5,9) Similarly, in two studies that focused on ctDNA at baseline, high ctDNA levels were associated with poor PFS (HR = 2.13; 95% CI: 1.43-3.17) and poor OS (HR = 1.80; 95% CI:0.95-

3.40) (7,11). The lower HR for survival outcomes at baseline for ctDNA compared to ccf-DNA observed in this meta-analysis could be due to the fewer number of studies available that assessed ctDNA at baseline; despite the fact that theoretically ctDNA is more specific representation of tumor activity as it is less likely to be affected by DNA from other cells/tissue.

Further, there are many challenges that need to be resolved to use ctDNA as a prognostic biomarker to monitor disease. Since, different genes and methods have been used by the researchers to measure the level of ctDNA. Hence, there is need to establish a common method and a similar set of genes or hot spot mutations for future prognosis, diagnosis, and analysis of ctDNA in DLBCL patients.

In our current meta-analysis, there are few limitations. The methodologies for ccf-DNA or ctDNA estimation of the included studies were not entirely uniform, which may contribute to heterogeneity in the results. The differences in patient characteristics could also be a source of variability. The sample sizes of included studies were small. There are no uniform threshold for ccf-DNA or ctDNA provided in the included studies. These limitations reduced the reliability of our findings. Our meta-analysis was limited to studies published in English.

CONCLUSION

The results of this meta-analysis strongly suggest that elevated levels of ccf-DNA or ctDNA are predictive of poor overall and progression free survival in patients with DLBCL. Additionally, these biomarkers have a strong potential to emerge as a valuable tool for monitoring disease status, such as MRD in DLBCL patients. As there are tumor heterogeneity in DLBCL patients

Highlights

1. Diffuse large B-cell lymphoma (DLBCL) is the most common type of non-Hodgkin lymphoma (NHL).
2. Relapses in DLBCL frequently arise from MRD that is not detectable by imaging techniques, highlighting the need for a sensitive and accurate biomarker to enhance the prediction of therapy response.
3. The use of ccf-DNA or ctDNA as a non-invasive biomarker has gained significant attention for the management of patients with DLBCL in the recent years.
4. Many studies have explored ccf-DNA or ctDNA as a prognostic biomarker for patients with DLBCL, but they are small, single center experience with inconsistent results.
5. This meta-analysis was conducted to evaluate the potential role of ccf-DNA or ctDNA in patients with DLBCL who were treated with combination chemotherapy (mainly R-CHOP or R-CHOP-like therapies)

6. This study could be a stepping stone toward utilizing ccfDNA or ctDNA as prognostic biomarker for DLBCL patients.

Acknowledgements

We are thankful to all the individuals who have participated in study and thus helped us produce this study's data.

Declarations of Interest statement

Availability of data and material: Yes

Competing interests: None

Funds: NA

Authors Contribution

AK: Conceptualization, Data extraction, analysis and writing

AK: Conceptualization, writing, supervision,

CS: Data extraction

AG: Software, Data analysis

PS: Review, editing, supervision

AJ: Review, editing, supervision

SS: Review, editing, supervision

AB: Review, editing, supervision

ARK: Review, editing, supervision

PM: Review, editing, supervision

GP: Conceptualization, Data extraction, writing, supervision

REFERENCES

1. Shi X, Liu X, Li X, Li Y, Lu D, Sun X, et al., Risk Stratification for Diffuse Large B-Cell Lymphoma by Integrating Interim Evaluation and International Prognostic Index: A Multicenter Retrospective Study. *Front Oncol.* 2021 Dec 16;11:754964. doi: 10.3389/fonc.2021.754964. PMID: 34976802; PMCID: PMC8716489.
2. Sogbe M, Aliseda D, Sangro P, de la Torre-Aláez M, Sangro B, Argemi J. Prognostic value of circulating tumor DNA in different cancer types detected by ultra-low-pass whole-genome sequencing. A systematic review and patient-level survival data meta-analysis. *Carcinogenesis.* 2024 Nov 16;bgae073. doi: 10.1093/carcin/bgae073. Epub ahead of print. PMID: 39549302.
3. Chen D, Guo J, Huang H, Tian L, Xie Y, Wu Q. Prognostic value of circulating tumor DNA in operable non-small cell lung cancer: a systematic review and reconstructed individual patient-data based meta-analysis. *BMC Med.* 2023 Nov 27;21(1):467. doi: 10.1186/s12916-023-03181-2. PMID: 38012727; PMCID: PMC10683311.
4. Kirchweiger P, Wundsam HV, Rumpold H. Circulating tumor DNA for diagnosis, prognosis and treatment of gastrointestinal malignancies. *World J Clin Oncol.* 2022 Jun 24;13(6):473-484. doi: 10.5306/wjco.v13.i6.473. PMID: 35949436; PMCID: PMC9244970.

5. Kristensen LS, Hansen JW, Kristensen SS, Tholstrup D, Harsløf LB, Pedersen OB, et al., Aberrant methylation of cell-free circulating DNA in plasma predicts poor outcome in diffuse large B cell lymphoma. *Clin Epigenetics*. 2016 Sep 7;8(1):95. doi: 10.1186/s13148-016-0261-y. PMID: 27610206; PMCID: PMC5015248.
6. Li M, Jia Y, Xu J, Cheng X, Xu C. Assessment of the circulating cell-free DNA marker association with diagnosis and prognostic prediction in patients with lymphoma: a single-center experience. *Ann Hematol*. 2017 Aug;96(8):1343-1351. doi: 10.1007/s00277-017-3043-5. Epub 2017 Jun 16. PMID: 28623396; PMCID: PMC5486778.
7. Kurtz DM, Scherer F, Jin MC, Soo J, Craig AFM, Esfahani MS, et al., Circulating Tumor DNA Measurements As Early Outcome Predictors in Diffuse Large B-Cell Lymphoma. *J Clin Oncol*. 2018 Oct 1;36(28):2845-2853. doi: 10.1200/JCO.2018.78.5246. Epub 2018 Aug 20. PMID: 30125215; PMCID: PMC6161832.
8. Chiu BC, Zhang Z, You Q, Zeng C, Stepniak E, Bracci PM, et al., Prognostic implications of 5-hydroxymethylcytosines from circulating cell-free DNA in diffuse large B-cell lymphoma. *Blood Adv*. 2019 Oct 8;3(19):2790-2799. doi: 10.1182/bloodadvances.2019000175. PMID: 31570490; PMCID: PMC6784517.
9. Hur JY, Kim YJ, Yoon SE, Son DS, Park WY, Kim SJ, et al., Plasma cell-free DNA is a prognostic biomarker for survival in patients with aggressive non-Hodgkin lymphomas. *Ann Hematol*. 2020 Jun;99(6):1293-1302. doi: 10.1007/s00277-020-04008-3. Epub 2020 Apr 15. PMID: 32296914.
10. Guan T, Zhang M, Liu X, Li J, Xin B, Ren Y, et al., Circulating tumor DNA mutation profile is associated with the prognosis and treatment response of Chinese patients with newly diagnosed diffuse large B-cell lymphoma. *Front Oncol*. 2022 Nov 17;12:1003957. doi: 10.3389/fonc.2022.1003957. PMID: 36465410; PMCID: PMC9713409.
11. Li M, Mi L, Wang C, Wang X, Zhu J, Qi F, et al., Clinical implications of circulating tumor DNA in predicting the outcome of diffuse large B cell lymphoma patients receiving first-line therapy. *BMC Med*. 2022 Oct 25;20(1):369. doi: 10.1186/s12916-022-02562-3. PMID: 36280874; PMCID: PMC9594942.
12. Alcoceba M, Stewart JP, García-Álvarez M, Díaz LG, Jiménez C, Medina A, et al., Liquid biopsy for molecular characterization of diffuse large B-cell lymphoma and early assessment of minimal residual disease. *Br J Haematol*. 2024 Jul;205(1):109-121. doi: 10.1111/bjh.19458. Epub 2024 May 29. PMID: 38811363.
13. Arzuaga-Mendez J, Prieto-Fernández E, Lopez-Lopez E, Martin-Guerrero I, García-Ruiz JC, García-Orad A. Cell-free DNA as a biomarker in diffuse large B-cell lymphoma: A systematic review. *Crit Rev Oncol Hematol*. 2019 Jul;139:7-15. doi: 10.1016/j.critrevonc.2019.04.013. Epub 2019 Apr 27. PMID: 31112884.
14. Yao L, Xu H, Wo J, Zhao M, Liu Z, Dong T, et al., Prognostic value of circulating tumor DNA in lymphoma: a meta-analysis. *Clin Exp Med*. 2022 Feb;22(1):1-7. doi: 10.1007/s10238-021-00718-8. Epub 2021 May 15. PMID: 33990849.
15. Tan X, Yan H, Chen L, Zhang Y, Sun C. Clinical Value of ctDNA in Hematological Malignancies (Lymphomas, Multiple Myeloma, Myelodysplastic Syndrome, and Leukemia): A Meta-Analysis. *Front Oncol*. 2021 Mar 4;11:632910. doi: 10.3389/fonc.2021.632910. PMID: 33747954; PMCID: PMC7970179.
16. Percival ME, Lai C, Estey E, Hourigan CS. Bone marrow evaluation for diagnosis and monitoring of acute myeloid leukemia. *Blood Rev*. 2017 Jul;31(4):185-192. doi: 10.1016/j.blre.2017.01.003. Epub 2017 Feb 2. PMID: 28190619; PMCID: PMC5513766.
17. Kim SH, Weigelt B. Circulating cell-free (cf)DNA analysis: Current technologies and applications in gynecologic cancer. *Gynecol Oncol Rep*. 2024 Jun 13;54:101431. doi: 10.1016/j.gore.2024.101431. PMID: 38947418; PMCID: PMC11211888.
18. Tabari E, Lovejoy AF, Lin H, Bolen CR, Lor Saelee S, Lefkowitz JP, et al., NGS-determined molecular markers and disease burden metrics from ctDNA correlate with PFS in previously untreated DLBCL. *Leuk Lymphoma*. 2024 May;65(5):618-628. doi: 10.1080/10428194.2024.2301924. Epub 2024 Feb 9. PMID: 38337191.
19. Kurtz DM, Green MR, Bratman SV, Scherer F, Liu CL, Kunder CA, et al., Noninvasive monitoring of diffuse large B-cell lymphoma by immunoglobulin high-throughput sequencing. *Blood*. 2015 Jun 11;125(24):3679-87. doi: 10.1182/blood-2015-03-635169. Epub 2015 Apr 17. PMID: 25887775; PMCID: PMC4463733.

20. Eskandari M, Manoochehrabadi S, Pashaiefar H, Zaimy MA, Ahmadvand M. Clinical significance of cell-free DNA as a prognostic biomarker in patients with diffuse large B-cell lymphoma. *Blood Res.* 2019 Jun;54(2):114-119. doi: 10.5045/br.2019.54.2.114. Epub 2019 Jun 25. PMID: 31309089; PMCID: PMC6614089.
21. Fu H, Shen J, Zhou H, Zhang F, Li H, Ma Z, et al., Mutation profiling of circulating tumor DNA identifies distinct mutation patterns in non-Hodgkin lymphoma. *Eur J Haematol.* 2022 Apr;108(4):298-309. doi: 10.1111/ejh.13736. Epub 2022 Jan 30. PMID: 34997652.
22. Figaredo G, Martín-Muñoz A, Barrio S, Parrilla L, Campos-Martín Y, Poza M, et al., Genetic Profiling of Cell-Free DNA in Liquid Biopsies: A Complementary Tool for the Diagnosis of B-Cell Lymphomas and the Surveillance of Measurable Residual Disease. *Cancers (Basel).* 2023 Aug 8;15(16):4022. doi: 10.3390/cancers15164022. PMID: 37627050; PMCID: PMC10452485.
23. Chen HY, Zhang WL, Zhang L, Yang P, Li F, Yang ZR, et al., 5-Hydroxymethylcytosine profiles of cfDNA are highly predictive of R-CHOP treatment response in diffuse large B cell lymphoma patients. *Clin Epigenetics.* 2021 Feb 11;13(1):33. doi: 10.1186/s13148-020-00973-8. PMID: 33573703; PMCID: PMC7879534.
24. Díaz-Chávez J, Gutiérrez-Hernández O, Taja-Chayeb L, Gutiérrez-Chavarría S, Avilés-Salas A, Candelaria M. ctDNA Is Useful to Detect Mutations at Codon 641 of Exon 16 of EZH2, a Biomarker for Relapse in Patients with Diffuse Large B-Cell Lymphoma. *Cancers (Basel).* 2022 Sep 24;14(19):4650. doi: 10.3390/cancers14194650. PMID: 36230571; PMCID: PMC9563768.
25. Le Goff E, Blanc-Durand P, Roulin L, Lafont C, Loyaux R, MBoumbae DL, et al., Baseline circulating tumour DNA and total metabolic tumour volume as early outcome predictors in aggressive large B-cell lymphoma. A real-world 112-patient cohort. *Br J Haematol.* 2023 Jul;202(1):54-64. doi: 10.1111/bjh.18809. Epub 2023 Apr 10. PMID: 37038217.
26. Li M, Ding N, Mi L, Shi Y, Du X, Yi Y, et al., Liquid biopsy in diffuse large B-cell lymphoma: utility in cell origin determination and survival prediction in Chinese patients. *Leuk Lymphoma.* 2022 Mar;63(3):608-617. doi: 10.1080/10428194.2021.1999441. Epub 2021 Nov 9. PMID: 34751093.
27. Roschewski M, Dunleavy K, Pittaluga S, Moorhead M, Pepin F, Kong K, et al., Circulating tumour DNA and CT monitoring in patients with untreated diffuse large B-cell lymphoma: a correlative biomarker study. *Lancet Oncol.* 2015 May;16(5):541-9. doi: 10.1016/S1470-2045(15)70106-3. Epub 2015 Apr 1. Erratum in: *Lancet Oncol.* 2015 May;16(5):e199. doi: 10.1016/S1470-2045(15)70217-2. PMID: 25842160; PMCID: PMC4460610.
28. Scherer F, Kurtz DM, Newman AM, Stehr H, Craig AF, Esfahani MS, et al., Distinct biological subtypes and patterns of genome evolution in lymphoma revealed by circulating tumor DNA. *Sci Transl Med.* 2016 Nov 9;8(364):364ra155. doi: 10.1126/scitranslmed.aai8545. PMID: 27831904; PMCID: PMC5490494.
29. Shirouchi Y, Mishima Y, Takayama T, Minowa S, Ishihara Y, Tamba M, et al., Serum cell-free DNA concentration as a possible prognostic marker in newly diagnosed diffuse large B-cell lymphoma. *Biomed Res.* 2022;43(4):99-106. doi: 10.2220/biomedres.43.99. PMID: 35989290.
30. Wedge E, Hansen JW, Garde C, Asmar F, Tholstrup D, Kristensen SS, et al., Global hypomethylation is an independent prognostic factor in diffuse large B cell lymphoma. *Am J Hematol.* 2017 Jul;92(7):689-694. doi: 10.1002/ajh.24751. Epub 2017 May 3. PMID: 28378885.
31. Al-Showbaki L, Wilson B, Tamimi F, Molto C, Mittal A, Cescon DW, et al., Changes in circulating tumor DNA and outcomes in solid tumors treated with immune checkpoint inhibitors: a systematic review. *J Immunother Cancer.* 2023 Feb;11(2):e005854.
32. Kruger DT, Jansen MPH, Konings IRHM, Dercksen WM, Jager A, Oulad Hadj J, et al., High ctDNA molecule numbers relate with poor outcome in advanced ER+, HER2- postmenopausal breast cancer patients treated with everolimus and exemestane. *Mol Oncol.* 2020 Mar;14(3):490-503.
33. Roosan MR, Mambetsariev I, Pharaon R, Fricke J, Husain H, Reckamp KL, et al., Usefulness of Circulating Tumor DNA in Identifying Somatic Mutations and Tracking Tumor Evolution in Patients With Non-small Cell Lung Cancer. *Chest.* 2021 Sep;160(3):1095-1107.