Circulating Cell-Free DNA and Colorectal Cancer: A Systematic Review

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Abstract: There is a strong demand for the identification of new biomarkers in colorectal cancer (CRC) diagnosis. Among all liquid biopsy analysts, cell-free circulating DNA (cfDNA) is probably the most promising tool with respect to the identification of minimal residual diseases, assessment of treatment response and prognosis, and identification of resistance mechanisms. Circulating cell-free tumor DNA (ctDNA) maintains the same genomic signatures that are present in the matching tumor tissue allowing for the quantitative and qualitative evaluation of mutation burdens in body fluids. Thus, ctDNA-based research represents a non-invasive method for cancer detection. Among the numerous possible applications, the diagnostic, predictive, and/or prognostic utility of ctDNA in CRC has attracted intense research during the last few years. In the present review, we will describe the different aspects related to cfDNA research and evidence from studies supporting its potential use in CRC diagnoses and the improvement of therapy efficacy. We believe that ctDNA-based research should be considered as key towards the introduction of personalized medicine and patient benefits.

Keywords: cell-free DNA; colorectal cancer; liquid biopsy

1. Introduction

The discovery of circulating acids (cell-free DNA-cfDNA, small and long non-coding RNAs) in body fluids attracted considerable attention in cancer research. The advantage of detection of cell-free DNA in plasma or serum is the potential to become a minimally invasive diagnostic and prognostic tool for colorectal cancer (CRC) patients [1]. Since early detection is a critical goal in cancer screenings, circulating nucleic acids represent a new class of biomarkers which can be used to improve the patients’ outcome.

Genetics represents a key role in predisposition to CRC and in its initiation, progression, and metastasis [2]. CRC provides a useful model for the study of other cancers—the various stages of tumor development can be observed. The well-defined sequence of events (from the aberrant crypt proliferation or hyperplasia via benign adenomas to carcinoma in situ and finally to metastatic carcinoma) testify the stepwise progression of this cancer, which often develops over many years [3]. The availability of biopsies is facilitated to determine that somatic mutations occur in several genes during the sequence of events. Apart from genetic events, smoking, diets, lifestyle, and other nongenetic factors are thought to have a strong impact on CRC risk.

Identifying predisposing germline mutations (heritable changes in the DNA occurring in the germ cells) together with acquired somatic mutations is important for the elucidation of the
carcinogenic process and in the design of diagnostic, therapeutic, and preventive strategies. Rapid advances are being achieved in the understanding of the molecular genetics and epigenetics of CRC. Accumulating evidence has presented CRC as a heterogeneous and complex disease. Molecularly and clinicopathologically, CRC is a very heterogeneous disease and different genetic features have been observed between colon cancer (CC) and rectal tumors [4]. These features implicate that the two diseases require different treatment strategies. A standard treatment of locally advanced rectal cancer consists of a multi-modal treatment with neoadjuvant chemoradiotherapy (therapy before surgery) followed by a total mesorectal excision of the tumor [5]. The common treatment for CC is surgical resection. If metastases are not present, the radical surgical resection of the primary tumor takes place. Because the surgery may not eliminate all cancer cells, adjuvant chemotherapy treatment is used to exterminate any tumor cells that may have been missed (micrometastasis). The adjuvant chemotherapy (applied after surgery) is used as a prevention method of cancer reappearance and is recommended for all patients with stage III CC without contraindications after curative resection. The medication 5-Fluorouracil (5-FU) has been the first choice of the adjuvant chemotherapy drug for CC for many years [6]. Usually, it is combined with leucovorin, which makes 5-FU more effective. Eventually, the combination of drugs called FOLFIRI (5-FU, leucovorin, and irinotecan) or FOLFOX (5-FU, leucovorin, and oxaliplatin) is applied [7].

CRC, a common neoplasia, ranks as the second most common type of cancer (11.5% worldwide), with nearly one million new cases diagnosed every year. An increase in the CRC incidence has been recorded all over Europe in the past decade, being particularly severe in central European regions [8]. Despite the continuously growing incidence, mortality rates gradually decrease in developed countries, probably due to the implementation of early screening [9] or the integration of new endoscopic and molecular techniques into clinical practice. Close collaboration between clinicians and scientists has expanded our understanding of the histo-clinico-pathological and molecular stages in CRC.

The prognosis for patients suffering from CRC is heavily dependent on the stage at diagnosis: a 5-year survival rate is up to 90% for stage I, but only <15% for advanced stages [6]. Over half of the cases are diagnosed at a higher stage of disease (III and IV). Treatment usually involves the complete primary tumor resection and appropriate chemotherapy. While the treatment can reduce the risk of relapse and increase a patient’s survival, it can also cause severe side effects and impair the quality of life. The differences in therapy response are mainly caused by a patient’s individual inherited genetic susceptibility affecting the response to the medication [10]. Current approaches to choose and implement chemotherapy regimens for CRC patients are primarily determined by tumor staging and histopathological examination. Developing diagnostic, prognostic, and predictive biomarkers based on the personal genetic background would greatly aid oncologists in the selection of the optimal type of drugs for each patient to improve their clinical outcome.

Significant effort is currently being employed to discriminate patients who will benefit from chemotherapy [11]. There is no current ‘gold standard’ to differentiate responders from non-responders. The analyses are usually performed by comparing the group of patients showing a certain response (from partial to complete) with a group of patients lacking any response.

Here, we will summarize the latest findings on cfDNA and evidence from studies supporting its potential use in CRC diagnoses and improvement of therapy efficacy.

1.1. Liquid Biopsy

There is a clear clinical need for novel diagnostic and molecular tools in oncology. Precision medicine focuses on the understanding of the precise relationship between the genes and phenotype, and the stratification of diseases into subtypes according to their underlying biological mechanisms. The heterogeneity of cancer indicates that the practice of using tissue biopsies for treatment decision making or disease monitoring has obvious weaknesses. Repeated monitoring of the tumor genome for treatment response assessment is a prerequisite for personalized therapy; unfortunately, this is nearly impossible because serial biopsies are usually a health burden for the patient due to the invasiveness
of the procedure [12]. Tumors are continuously in evolution, and even if several biopsies are obtained, these are limited both spatially and temporally. Moreover, in several patients, it is impossible to obtain biopsy samples and for sufficient material of adequate quality for genomic profiling, we need to be aware that reported failure rates range between 10–30% of cases [13]. It is important to keep in mind that almost all tumors treated with any therapy acquire resistance because of tumor heterogeneity, clonal evolution, and/or selection [14]. Usually, the treatment decision is based on the result from single tumor biopsy and many substantial lesions might by simply overlooked [12]. As stated above, in most cases, it is impossible to obtain a biopsy sample and no information about the genetic background of the developed metastasis is available. Moreover, during the evolution and progression of cancer, metastases might lose aberrations present in the primary lesion and this leads to differences between the primary tumor and the metastases.

With the improvements in research, it is now well known that various cells within a tumor constantly release many biomolecules such as DNA, RNA, and proteins into body fluids. These biomolecules can be either cell-free or bound to proteins or lipids, or capsulated in extracellular vesicles. Because circulating cell-free DNA (cfDNA) is a surrogate for the tumor genome, it is often referred to as a liquid biopsy [12]. Biomarkers in body fluids are thus of tremendous potential in translational research. A liquid biopsy minimizes invasive approaches to sampling cancer cells and their molecular signatures. Moreover, a liquid biopsy is a representative sample of the entire tumor, which overcomes tumor heterogeneity which is the difficult to be captured by tissue biopsy ([15], Figure 1) and represents a reliable source of diagnostic DNA and could thus replace the use of tumor tissue in diagnostic settings [16]. The fecal occult blood testing (FOBT) and colonoscopy are routinely used for the early detection of CRC. However, the sensitivity for detecting adenomas with FOBT is very low [17]. Moreover, the compliance for colonoscopy is quite low because it is time consuming, unpleasant, painful, and involving some risk [18]. A CRC screening test that accurately detects advanced adenomas with a high potential of malignant progression is needed [19]. Recently, the multitarget stool DNA test or the blood-based Septin 9 DNA methylation test have been launched in clinical practice. A liquid biopsy analysis may also exhibit a great diagnostic potential in CRC for monitoring resistance development to treatment. These new diagnostic tools and the definition of molecular biomarkers in CRC will improve early detection and targeted therapy of CRC. We believe that the combination of FOBT and a liquid biopsy analysis may improve the performance of CRC screenings.

![Figure 1](image_url). A liquid biopsy for solid tumors. A literature scan of the last few years revealed an enormous increase in liquid biopsy-based analyses [20].
The main differences between the conventional analysis of tumor tissue and a liquid biopsy are the fact that tumor tissue is very genetically heterogeneous, and tumor cells are, therefore, not all genetically identical to one another. The DNA, mRNA, or microRNA profiling of tumor tissue only provides the analysis of the predominant tumor cells. Modern high-throughput techniques enable the identification of several predominant tumor cells but not of all tumor cells unless every tumor cell is analyzed. Owing to this, several tumor cells may not have been captured and tested. However, this can be omitted by analyzing the liquid biopsy, albeit with a lower tumor fraction in the plasma, due to which the sensitivity of ctDNA detection decreases. Nevertheless, tissue and liquid biopsies investigate different parameters and, therefore, do not deliver identical results.

A ctDNA/cfDNA analysis has their own advantages and disadvantages and their results can supplement each other (Table 1).

Table 1. The advantages and disadvantages of a liquid biopsy.

| Advantages                                      | Disadvantages                                      |
|------------------------------------------------|---------------------------------------------------|
| Cost-effective                                  | Lack of standard operating protocol               |
| Non-invasive                                    | Released by both healthy and tumor cells          |
| Rapid                                           | Requirement of sensitive and specific methods     |
| Comprehensive tumor profile                     | False-positive and false-negative results         |
| Minimal pain and risk                           | Personnel microenvironment may influence the released of cfDNA amount |
| Serial assessments                               | Not-standardized cfDNA/ctDNA concentration as a cancer biomarker |
| Directly able to assess for specific mutations  |                                                   |
| Present in many biological fluids              |                                                   |
| Potential to evaluate prognosis, recurrence, response to therapy |   |
| Detection of minimal residual disease           |                                                   |
| Assessment of cancer high-risk populations      |                                                   |
| Enabling of early cancer diagnosis              |                                                   |
| Evaluation of tumor heterogeneity               |                                                   |

1.2. Cell-Free DNA

Circulating tumor DNA (ctDNA) is fragmented DNA derived from a tumor into the bloodstream and is not associated with cells. Very often the ctDNA term is confused with cfDNA, which characterizes DNA freely circulating in the bloodstream and is not necessarily of tumor origin. Because ctDNA reflects the entire tumor genome, it has earned an attraction for its potential clinical utility as a ‘liquid biopsies’. The fraction of ctDNA can range between 0.01% to 90% [21–23].

For the first time, cfDNA was discovered in the blood of healthy individuals by Mandel and Métais [24]. Higher cfDNA levels were first reported in the serum of patients with various types of cancer by Leon and Shapiro in 1977 and thus proved the diagnostic features of cfDNA in cancer patients. Similarly, metastatic cancer patients had a higher level of cfDNA than non-metastatic patients. The authors also observed that after radiation therapy, cfDNA levels decreased; albeit with constant or increasing levels of cfDNA being associated with a worse prognosis or relapse. Leon et al. hypothesized that cfDNA in serum could represent a promising biomarker for evaluating therapy response [25]. The following studies aimed to prove that cfDNA characterizes neoplastic features of the tumor. In 1989, Stroun et al. [26] suggested that part of the cfDNA in the plasma of cancer patients originates from tumor cells. Tumor-specific aberrations, like mutations in oncogenes and tumor suppressor genes [27], microsatellite instability (MSI) [28], and DNA methylation [29] were identified in cfDNA and studies confirmed the release of cfDNA into the circulation by tumors.
Recent reports have shown the possibility of reconstructing the genome of tumors from ctDNA as ctDNA represents a potential surrogate of the entire genome [30,31]. This means that genetic and epigenetic signatures in ctDNA correspond to those in the primary tumor and may reveal the tumor-specific (epi)-genetic spectrum [32]. This implies ctDNA is directly released from tumors. Several health conditions may be associated with increasing levels of cfDNA such as inflammation, tissue trauma, autoimmune diseases, or cancer [33,34]. Concerning cancer and cfDNA studies, most of the studies confirmed that patients with cancer have higher levels of cfDNA than patients with benign diseases or healthy individuals [35–38].

CfDNA could be found in serum, plasma, and other body fluids like urine or saliva [39,40] but the mechanisms of release into the bloodstream are not completely understood. In general, cfDNA may be derived from a primary tumor, metastatic lesions, or circulating tumor cells (CTCs) [41]. There are two ways of release—passive and active. Passive release means by necrotic and apoptotic cells and active secretion can be mediated by nucleated cells such as lymphocytes. Regardless, the major source of cfDNA in plasma is from the apoptotic or necrotic cells [42]. A few authors suggest that malignant tumors exhibit a higher degree of necrosis which corresponds to an increase in ctDNA levels. Diehl et al. [43] suggested that DNA fragments in the circulation are derived from necrotic tumor cells absorbed by macrophages. However, Leon et al. [25] showed that radiation therapy in patients induces cell necrosis and observed a 90% decrease in cfDNA levels. In addition, several studies stated that cfDNA is derived from active cellular secretion, for example, macrovesicles such as exosomes [44,45]. Recently, it was hypothesized that all cells that are alive actively release DNA into the circulation [12]. Therefore, both apoptosis and necrosis, alongside with active secretion play important role in the cfDNA presence in liquid biopsies.

cfDNA is typically fragmented into 180 bp fragments, corresponding to the length of DNA wrapped around a nucleosome and protected from degradation [12,46]. Another peak representing the multimers of nucleosomes in a minor fraction of the total can be observed [12]. A peak at 166 bp [47,48] corresponds to the length of DNA wrapped around a nucleosome (~147 bp) with DNA associated histone H1. Depending on the size of the fragment of cfDNA, the source of release can be identified. CfDNA fragments produced by apoptotic cells are about 200 bp long and contrarily cfDNA released from necrotic cells has a variable shorter length [42] and may thus increase the proportion of shorter fragments and lead to low integrity [49]. The ratio of longer to shorter DNA fragments is named the cfDNA integrity number and is determined as a ratio of two qPCR products of different lengths. Umetani et al. developed a method for measuring the integrity of cfDNA by qPCR for ALU repeats (247 bp ALU vs. 115 bp ALU) [50]. The reason for choosing these sequences is the argument that ALU repeats are the most abundant repeated sequences in the human genome [51]. CtDNA molecules are also shorter than non-mutated cfDNA molecules in plasma [48,52–55]. Longer cfDNA fragments (>1000 bp) were observed in healthy volunteers [56]. The authors hypothesized that the release into circulation in healthy individuals may be associated with exosomes [44,56–58] and in cancer patients by necrosis from tumor cells [42].

The average concentration of cfDNA in the blood serum of healthy volunteers was 13 ng/mL, whereas in cancer patients the mean level was 180 ng/mL [25]. The amount of circulating DNA, whether tumoral or not, is limiting: a range between 1 to 10 ng per ml plasma [59,60], which correspond to 1500 to 3000 copies of the haploid genome [61]. A substantial variation in ctDNA levels may arise from an interindividual difference. Poor tumor vascularization could hamper ctDNA release into the bloodstream or, on the other hand, could support ctDNA release by producing hypoxia and cell death [58]. When the plasma cfDNA and serum cfDNA concentration was compared, it was found out that the serum concentration is 3–24 higher than in plasma [52,63]. Jung et al. confirmed these differences and related it to the time delay and storage temperature of blood before centrifugation. Higher levels of cfDNA in the serum can also be caused by contamination by cells during the clotting process. For this and other reasons for the analysis of tumor-specific DNA, it is recommended to use the plasma as a source of cfDNA due to lower concentrations of awild-type DNA background [64].
Many researchers choose plasma for their analysis because it is less likely to be contaminated by leucocytes than serum [65]. It is important to remember that most of cfDNA originate from normal cells, mostly from the hematopoietic compartment, and ctDNA generally represents only a minor fraction of it, possibly 0.1% of the total amount, or even less [66,67].

cfDNA is considered a good biomarker and can be characterized in disease monitoring by two options—quantitative and/or qualitative changes. Quantitative changes include differences in the concentration of cfDNA and qualitative changes are represented by gene mutations, loss of heterozygosity, DNA copy number variations, methylation, microsatellite instability (MSI), etc.

As ctDNA represents only a very small proportion of cfDNA, very sensitive and reliable detection methods are required. Detection methods can be divided into two groups: (i) the targeted approach that allows for the detection of specific alterations, and (ii) the untargeted approach that allows for the identification of events a priori, for example, whole exome or genome sequencing. Levels of cfDNA/ctDNA are measured mainly by real-time PCR (RT-PCR) [68]. Digital PCR (dPCR), RT-PCR [27,69] or sequencing methods [70], beads Emulsion Amplification and Magnetics (BEAMing) are used for the detection of point mutations. Sequencing methods can identify structural rearrangements, chromosomal copy-number changes, and structural alternations of ctDNA in circulation [71,72]. Different cfDNA/ctDNA detection techniques with their pros and cons are summarized in Table 2.

Table 2. A summary of ctDNA detection techniques (adopted from References [73,74]).

| Technique | Limit of Detection | Type of Alteration Detection | Advantages | Disadvantages |
|-----------|-------------------|-----------------------------|------------|---------------|
| PCR based approaches (COLD-PCR, PNAs-LNA, ARMS, etc.) | 0.1–1% | SNV, indels | Low cost; Easy to perform | Low sensitivity; A limited number of studied genes at a time; Genes need to be pre-determined |
| Digital PCR (dPCR and BEAMing) | 0.05% or less | SNV, indels, CNV | High sensitivity and specificity; Reasonable cost; Easy to perform | A limited number of studied genes at a time; Genes need to be pre-determined |
| NGS (Deep sequencing, TAM-seq, Safe-Seqs, CAPP-Seq, cSMART, digital sequencing) | 0.01–2% | SNV, indels, CNV, rearrangements | Allows more genes to be analyzed at a time | Wide range of sensitivity depending on the NGS platform used (PCR amplicon strategies are more sensitive and less expensive than whole genome or exome sequencing); Higher cost |
| SERS-nanotags | 0.01% | SNV | Reduced susceptibility to photobleaching; Bandwidths are significantly narrower | Raman signal deterioration upon prolonged laser illumination |
| UltraSEEK | 0.01% | SNV, indels | Low cost and low DNA input | Lower sensitivity |

cfDNA: circulating tumor DNA, COLD-PCR: co-amplification at lower denaturation temperature; PNAs-LNA: peptide nucleic acid-locked nucleic acid; ARMs: amplification refractory mutations system; dPCR: droplet digital PCR; BEAMing: bead emulsion amplification and magnetics; NGS: next-generation sequencing; TAM-Seq: tagged-amplicion deep sequencing; Safe-Seq: safe sequencing system; CAPP-Seq: cancer personalized profiling by deep sequencing; cSMART: circulating single molecule amplification and re-sequencing technology; SERS: surface-enhanced Raman spectroscopy; UltraSEEK: high-throughput multiplex ultrasensitive mutation detection; SNV: single nucleotide variation; CNV: copy number variation.
1.3. Preanalytical Considerations

The preparation of cfDNA for analysis is easy to implement. For the isolation of cfDNA, about 5–10 mL of blood collected in tubes with anticoagulants is required. CfDNA could be processed from serum as well but with a lower priority due to the lysed cellular DNA that may have an impact on relative levels of cfDNA. In the blood, cfDNA has a limited stability because of the DNase activity and therefore the preparation of cfDNA should not overstep three hours after blood collection [41]. Cell lysis needs to be avoided in order to prevent the release of a large amount of non-mutant DNA, potentially leading to a false negative result [67].

The lack of consistency between several protocols for sample handling and methodologies used for cfDNA analyses represent the major obstacles in translating cfDNA based research into clinical practice. Therefore, Messaoudi et al. [75] and recently Nikolaev [67] determined the optimal preanalytical protocols for cfDNA analyses:

- Plasma is a better source of cfDNA than serum since it avoids blood cell genomic DNA contamination
- EDTA or cell-free DNA™ collection tubes prevent blood cell lysis by keeping tubes at 4 °C
- Blood must be processed within a maximum of 4 h following blood drawing to preserve cfDNA concentration and fragmentation
- To ensure any absence of cells in plasma, first, centrifugation is recommended at 1200–1600 g for 10 min and, second, microcentrifugation at 16,000 g for 10 min, (the second step can be indifferentely realized before or even after storage of plasma samples)
- Plasma samples must be stored at −80 °C for up to nine months (samples are sensitive to freeze-thaw cycles)
- cfDNA extracts may tolerate a maximum of three freeze-thaw cycles and storage at −20 °C for up to three months

2. Results

2.1. Cell-Free DNA as a Physiological Mobile Genetic Element

Mobile genetic elements play an important role in shaping biotic genomes and bringing about evolutionary transformations [76]. Recently it was observed that cfDNA, after entering healthy cells, integrated into their genomes and led to DNA damage [77].

Genomic integration of cfDNA might then lead to DNA rearrangements, translocations, and deletions [78] and ultimately induce the aging of cells [79] or may activate chemoresistance.

A process called genometastasis, where the cfDNA in the plasma may participate in tumorigenesis and the development of metastases via transfection-like uptake of nucleic acids by susceptible cells was recently introduced. García-Olmo [80] cultured NIH-3T3 mouse cells with plasma samples from CRC patients bearing KRAS mutation (codon 12 in exon 1), as well as with plasma from healthy individuals. The authors detected mutated human KRAS sequences in cultures of NIH-3T3 cells after the start of incubation and these sequences were still detectable at the end of the experiment, even ~3 weeks after the removal of human plasma from the culture medium. NIH-3T3 cells that were cultured with plasma from healthy individuals were negative. NIH-3T3 cells (p16-deficient cells) were able to stably incorporate foreign DNA during a simple incubation with plasma from cancer patients. Another study demonstrated the integration of ctDNA into nuclear DNA of leukemic cells and suggested that this process may occur through non-homologous end-joining [81]. This all can conclude that cfDNA, far from being an inert molecule, has some biological functions of their own that are deleterious to healthy cells of the body.
2.2. Cell-Free DNA and Colorectal Cancer

Analysis of ctDNA is a promising new tool in oncology. ctDNA mutational content can provide invaluable information on the genetic background of a tumor, and assist oncologist in deciding on therapy, or in following the residual disease (Figure 2, Table 3).

**Figure 2.** A liquid biopsy to monitor the therapy response and resistance. A hypothetical scenario of the developing chemotherapy resistance of colorectal cancer patients: first-line therapy is based on the primary tumor and relevant changes in the metastasis might be overlooked, therefore, leading to primary resistance. After switching to second-line therapy, secondary resistance may arise. Genetic changes of the resistant clones can be monitor using liquid biopsy and, therefore, the resistance mechanism might be recognized before disease progression (adapted from Heitzer et al. [12]).
### Table 3. An overview of the studies investigating cfDNA/ctDNA in relation to CRC diagnosis, therapy outcome, and prognosis.

| References        | Patients | Controls | Origin of the Study | Source of cfDNA/ctDNA | Abnormalities | Methodology         | Target    | Clinical Relevance |
|-------------------|----------|----------|---------------------|-----------------------|---------------|---------------------|-----------|-------------------|
| Leon et al. [25]  | 9        | 55       | USA                 | Serum                  | Concentration  | Diagnostic          |           |                   |
| Anker et al. [82] | 14       |          | Switzerland         | Plasma                 | Mutation       | PCR                 | KRAS      | Diagnostic         |
| de Kok et al. [83]| 14       |          | Netherlands         | Serum                  | Mutation       | PCR                 | KRAS      | Diagnostic         |
| Kopreski et al. [84]| 31      | 28       | USA                 | Plasma Serum           | Mutation       | PCR                 | KRAS      | Diagnostic         |
| Hibi et al. [85]  | 44       |          | USA                 | Serum                  | Mutation       | PCR                 | KRAS      | Diagnostic         |
| de Kok et al. [83]|          |          | Netherlands         | Serum                  | Mutation       | PCR                 | KRAS      | Diagnostic         |
| Ryan et al. [87]  | 94       |          | Ireland             | Serum                  | Mutation       | PCR sequencing      | KRAS2     | Diagnostic         |
| Wang et al. [27]  | 104      | 50       | Taiwan              | Serum                  | Mutation Concentration | PCR-SSCP    | APC       | Diagnostic         |
| Leung et al. [88] | 49       | 41       | Hong Kong           | Serum                  | Methylation    | Methylight          | APC       | Diagnostic         |
| Lindforss et al. [89]| 25      |          | Sweden              | Plasma                 | Mutation       | PCR                 | KRAS      | Prognostic         |
| Bazan et al. [90] | 66       |          | Italy               | Plasma                 | Mutation Methylation | PCR         | KRAS      | Prognostic         |
| Flamini et al. [91]| 75      | 75       | Italy               | Serum                  | Concentration  | qPCR                |           | Diagnostic         |
| Frattini et al. [92]| 70      | 20       | Italy               | Plasma                 | Concentration  | qPCR                |           | Diagnostic         |
| Trevisiol et al. [93]| 86      |          | Italy               | Serum                  | Mutation       | qPCR                | KRAS2     | Diagnostic         |
| Wallner et al. [94]| 38      | 20       | Germany             | Serum                  | Methylation    | MS-PCR              | HPP1/TP6F | Prediction         |
| Boni et al. [95]  | 67       | 67       | Italy               | Plasma                 | Concentration  | qPCR                |           | Diagnostic         |
| Nakayama et al. [96]| 94      |          | Japan               | Serum                  | Methylation    | MS-PCR              | p16       | Diagnostic         |
| References                        | Patients | Controls | Origin of the Study | Source of cfDNA/ctDNA | Abnormalities | Methodology | Target          | Clinical Relevance     |
|----------------------------------|----------|----------|---------------------|-----------------------|---------------|-------------|-----------------|------------------------|
| Diehl et al. [97]                | 18       | -        | USA                 | Plasma                | Concentration | qPCR        |                 | Diagnostic Prognostic  |
| Frattini et al. [98]             | 70       | 20       | Italy               | Plasma                | Concentration | qPCR, MS-PCR, ME-PCR | KRAS, p16INK4a         | Diagnostic Predictive  |
| Lofton-Day et al. [99]           | 133      | 179      | Germany             | Plasma                | Methylation   | MS-PCR      | TMEFF2, NGFR, SEPT9 | Diagnostic             |
| Schwarzenbach et al. [100]       | 55       | 20       | Germany             | Serum                 | Concentration | qPCR        |                 | Diagnostic             |
| Su et al. [39]                   | 20       | -        | USA                 | Serum, Plasma, Urine | Concentration | PCR         | KRAS            | Diagnostic             |
| Zitt et al. [101]                | 26       | -        | Austria             | Plasma                | Concentration | qPCR        |                 | Prognostic Predictive  |
| DeVos et al. [102]               | 97       | 172      | Germany             | Plasma                | Methylation   | MS-PCR      | SEPT9           | Diagnostic             |
| Herbst et al. [103]              | 106      | -        | Germany             | Serum                 | Methylation   | Methylight  | HLF, HPP1, TPEF | Prognostic             |
| Agostini et al. [104]            | 67       | 35       | Italy               | Plasma                | Concentration | qPCR        |                 | Predictive             |
| Herbst et al. [103]              | 106      | -        | Germany             | Plasma                | Methylation   | MS-PCR      | HPP1, TPEF, HLF, NEUROG1 | Diagnostic             |
| Diaz et al. [105]                | 28       | -        | USA                 | Serum                 | Mutation      | PCR         | KRAS            | Diagnostic Predictive  |
| Morgan et al. [106]              | 71       | -        | UK                  | Plasma, Serum         | Mutation      | qPCR        | KRAS            | Diagnostic             |
| Phillip et al. [107]             | 311      | -        | Germany             | Serum                 | Methylation   | MS-PCR      | HLF, HPP1       | Prognostic             |
| Spindler et al. [108]            | 108      | -        | Denmark             | Plasma                | Concentration | qPCR        | KRAS            | Prognostic Predictive  |
Table 3. Cont.

| References          | Patients | Controls | Origin of the Study | Source of cfDNA/ctDNA | Abnormalities | Methodology | Target | Clinical Relevance |
|---------------------|----------|----------|---------------------|-----------------------|---------------|-------------|--------|--------------------|
| Bai et al. [109]    | 106      | -        | China               | Plasma                | Mutation      | PCR         | KRAS   | Diagnostic         |
|                     |          |          |                     |                       |               |             |        | Prognostic         |
| Cassinotti et al. [110] | 223      | -        | Italy               | Plasma                | Concentration | qPCR        |        | Prognostic         |
| Lee et al. [111]    | 101      | 96       | Korea               | Plasma                | Methylation   | PCR         | Septin9| Diagnostic         |
| Spindler et al. [112] | 211      | -        | Denmark             | Plasma                | Mutation      | qPCR        | KRAS   | Predictive         |
|                     |          |          |                     |                       |               |             | BRAF   | Prognostic         |
| Sun et al. [113]    | 34       | 10       | China               | Plasma                | Concentration | qPCR, MS-PCR, PCR-RFLP | MGMT   | Diagnostic         |
|                     |          |          |                     |                       | Methylation   |             | KRAS   | Prognostic         |
| Bettegowda et al. [21] | 24       | -        | USA                 | Plasma                | Concentration | PCR         | KRAS   | Diagnosis          |
| Kuo et al. [16]     | 52       | -        | Taiwan              | Plasma                | Mutation      | PCR         | KRAS   | Predictive         |
| Lin et al. [114]    | 133      | -        | Taiwan              | Plasma                | Mutation      | qPCR        | 74 genes| Prognostic         |
| Mohan et al. [115]  | 10       | -        | Austria             | Plasma                | Mutation      | WGS         | KRAS   | Diagnostic         |
|                     |          |          |                     |                       |               |             | BRAF   | Predictive         |
|                     |          |          |                     |                       |               |             | PIK3CA | Prognostic         |
|                     |          |          |                     |                       |               |             | EGFR   | Prognostic         |
| Perrone et al. [116] | 170      | -        | Italy               | Plasma                | Mutation      | ME-PCR, qPCR | KRAS   | Diagnostic         |
|                     |          |          |                     |                       | Concentration |             |        |                    |
| Spindler et al. [117] | 108      | -        | Denmark             | Plasma                | Mutation      | PCR         | KRAS   | Predictive         |
|                     |          |          |                     |                       | Concentration |             | BRAF   | Prognostic         |
| Spindler et al. [118] | 100      | 100      | Denmark             | Plasma                | Concentration | PCR         | KRAS   | Diagnostic         |
|                     |          |          |                     |                       | Mutation      |             |        | Predictive         |
| Tham et al. [119]   | 150      | -        | Singapore           | Serum                 | Methylation   | MS-PCR      | TACI   | Prognostic         |
|                     |          |          |                     |                       |               |             | Septin9|                  |
|                     |          |          |                     |                       |               |             | NELL1  |                  |
| Thierry et al. [23] | 106      | 29       | France              | Plasma                | Mutation      | qPCR        | KRAS   | Diagnostic         |
|                     |          |          |                     |                       |               |             | BRAF   | Predictive         |
| Xu et al. [120]     | 242      | -        | China               | Plasma                | Mutation      | PCR         | KRAS   | Prognostic         |
| References                        | Patients | Controls | Origin of the Study | Source of cfDNA/ctDNA | Abnormalities | Methodology | Target | Clinical Relevance |
|----------------------------------|----------|----------|---------------------|-----------------------|---------------|-------------|--------|-------------------|
| Carpinetti et al. [121]          | 4        | -        | Brazil              | Plasma                | Chromosomal Rearrangements | SOLiD        |        | Predictive         |
| Lin et al. [122]                 | 353      | -        | Taiwan              | Plasma                | Methylation    | Methylation array | >450,000 CpG sites | Diagnostic |
| Reinert et al. [123]             | 118      | -        | Denmark             | Plasma                | Concentration  | ddPCR       |        | Diagnostic         |
| Sefrioui et al. [124]            | 34       | -        | France              | Plasma                | Mutation       | dPCR        | KRAS   | Diagnostic         |
| Siravegna et al. [125]           | 100      | -        | Italy               | Plasma                | Mutation       | PCR         | KRAS   | Prognostic         |
| Spindler et al. [126]            | 229      | 100      | Denmark             | Plasma                | Mutation       | qPCR        | KRAS   | Diagnostic         |
| Liu et al. [127]                 | 165      | -        | Singapore           | Serum                 | Methylation    | MS-PCR      | SST    | Prognostic         |
| Matthaios et al. [128]           | 155      | -        | Greece              | Plasma                | Methylation    | MS-PCR      | APC    | Prognostic         |
| El Messaoudi et al. [129]        | 97       | -        | Francie             | Plasma                | Mutation       | qPCR        | KRAS   | Diagnostic         |
| Tie et al. [130]                 | 230      | -        | Australia           | Plasma                | Mutation       | PCR         | APC    | Prognostic         |
| Agah et al. [131]                | 74       | -        | Iran                | Plasma                | Concentration  | qPCR        | TP53   | Diagnostic         |
| Bhang et al. [132]               | 30       | 17       | Austria             | Plasma                | Concentration  | qPCR        |        | Diagnostic         |
| Herbst et al. [133]              | 467      | -        | Germany             | Plasma                | Methylation    | MS-PCR      | HPP1   | Predictive         |
| Kloten et al. [134]              | 50       | 8        | Germany             | Plasma                | Concentration  | qPCR mutation | KRAS   | Diagnostic         |
| Li et al. [135]                  | 80       | 35       | USA                 | Serum Plasmap         | Concentration Cnvs | WGS        |        | Diagnostic         |
| Pereira et al. [136]             | 128      | -        | USA                 | Plasma                | Mutation       | sequencing  |        | Diagnostic         |
| Yamauchi et al. [137]            | 21       | -        | Japan               | Plasma                | Mutation       | sequencing  |        | Predictive         |
| Liu et al. [138]                 | 27       | -        | USA                 | Plasma                | Methylation    | Infinium HM450 array |        | Diagnostic         |
| References               | Patients | Controls | Source of cfDNA/ctDNA | Abnormalities | Methodology | Target | Clinical Relevance |
|--------------------------|----------|----------|-----------------------|---------------|-------------|--------|--------------------|
| Takayama et al. [139]    | 85       | -        | Japan                 | Plasma        | Concentration Mutations | dPCR   | KRAS               |
| Toledo et al. [140]      | 1        | -        | Spain                 | Plasma        | Whole Exome Sequencing  | sequencing | Predicitive   |
| Schou et al. [141]       | 123      | -        | Denmark               | Plasma        | Concentration fluorescence | dPCR   | KRAS             |
| Sclafani et al. [142]    | 51       | -        | Clinical Trial        | Plasma        | Mutation               | ddPCR  | KRAS, BRAF        |
| Boysen et al. [143]      | 273      | 94       | Denmark, Norway, Sweden | Plasma        | Concentration          | ddPCR, DFA | Diagnostic  |
| Myint et al. [144]       | 131      | 37       | UK                    | Plasma, Stool | Concentration Mutations | qPCR   | KRAS, BRAF        |
| Demuth et al. [145]      | 28       | -        | Denmark               | Plasma        | Mutation               | ddPCR  | KRAS             |
| Rokni et al. [146]       | 50       | -        | Iran                  | Plasma        | Methylation            | ddPCR, High methylation resolution PCR | BMP3 | Prognostic |
| Fu et al. [147]          | 98 CRC, 101 adenomas, 76 nCRC | 253 | China | Plasma | Methylation | MS-PCR | SEPT9 | Prognostic |
| Molpahia et al. [148]    | 24       | 25       | USA                   | Plasma        | Cnvs                    | sequencing | Diagnostic |
| Gallardo-Gómez et al. [149] | 20 CRC, 20 adenomas | | Spain | Serum | Methylation | microarray | APC, FOXA1, MGMT, RARβ2, RASSF1A, SCGB3AI, SEPT9, SHOX2, SOX17 | Prognostic |
| Nunes et al. [150]       | 72       | 103      | Portugal              | Plasma        | Methylation            | MS-PCR |                     |
| References            | Patients | Controls | Origin of the Study | Source of cfDNA/ctDNA | Abnormalities | Methodology | Target          | Clinical Relevance |
|-----------------------|----------|----------|---------------------|-----------------------|---------------|-------------|------------------|-------------------|
| Song et al. [151]     | 150      | -        | China               | Urine                 | Concentration | ddPCR       |                  | Predictive        |
| Yang et al. [152]     | 47       | -        | China               | Plasma                | Mutation      | sequencing  | 37 genes        | Prognostic         |
| Suehiro et al. [153]  | 113      | 25       | Japan               | Serum                 | Methylation   | ddPCR       | TWIST1           | Diagnostic         |
| Sun et al. [154]      | 11       | -        | China               | Plasma                | Mutation      | sequencing  | 85 genes        | Prognostic         |
| Thomsen et al. [155]  | 138      | -        | Denmark             | Plasma                | Concentration | ddPCR       | RAS/RAF          | Prognostic         |
| Furuki et al. [156]   | 22       | -        | Japan               | Serum                 | Mutation      | sequencing  | TP53, KRAS, APC, PIK3CA, BRAF, FBXW7, NRAS | Diagnostic         |
| Klein-Scory et al. [157]| 3        | -        | Germany             | Plasma                | Mutation      | BEAMing     | BRAF, PIK3CA   | Predictive        |
| Schøler et al. [158]  | 45       | -        | Denmark             | Plasma                | Concentration | WGS         |                  | Prognostic         |
| Vandeputte et al. [159]| 20      | -        | Belgium             | Plasma                | Concentration | ddPCR       |                  | Predictive        |

cfDNA: circulating cell-free DNA, ctDNA: circulating tumor DNA, CRC: colorectal cancer, ddPCR: droplet digital PCR; DFA: direct fluorescent assay; dPCR: digital PCR; MS-PCR: methylation specific PCR; qPCR: quantitative PCR; RT-PCR: real-time PCR; SOLID: Sequencing by Oligonucleotide Ligation and Detection; SSCP: single strand conformation PCR; WGS: whole genome sequencing.
2.2.1. Cell-Free DNA as a Diagnostic Biomarker in Colorectal Cancer

The average concentration of serum cfDNA from CRC patients was 5-times higher than that in the serum of healthy controls, while in plasma, it was 25–50 times higher in CRC patients than in the plasma of healthy controls [91,92,95,126]. Several authors analyzed the cfDNA concentration differences in plasma between rectal and colon cancer patients. Colon cancer patients evinced higher cfDNA concentration than patients with rectal cancer (RC) (colon: 500 ng/mL, rectum: 250 ng/mL in plasma) [98], while Cassinotti et al. observed a higher concentration of cfDNA in patients with RC [110].

Activating mutations in the KRAS gene are predictors of poor response in patients with metastatic CRC receiving anti-EGFR (epidermal growth factor receptor) antibody-based therapy. Therefore, these patients are routinely tested for the presence of KRAS mutations before receiving biological therapy [160,161]. Numerous studies tried to prove that KRAS mutation can be detected in cfDNA and may serve as a diagnostic tool. Already in 1997, Anker et al. [82] postulated that the genetic analysis of plasma DNA may have clinical applications in the future. In their study, KRAS abnormalities were analyzed in 14 tumors and plasma from CRC patients. Similarly, de Kok and Kopreski, in 1997 [83,84], analyzed the serum of CRC patients for KRAS mutations. Back in 1992, Sidransky et al. [162] reported that KRAS mutations in tumors from CRC patients and stool samples were identical. The possibility of detecting mutations in body fluids like plasma and serum was analyzed in many other studies [86,115,126]. Wang et al. detected KRAS mutations in tumor tissue and evaluated the presence of these mutations in serum samples. Their results showed that about 45% of CRC patients with KRAS mutations in tumor tissues evinced these mutations in cfDNA. Interestingly, KRAS mutations were not detected in the cfDNA of the healthy control group [27]. The concordance between BRAF and KRAS mutation analysis in tumor tissue and matched plasma was analyzed further. The KRAS mutation analysis showed a 96% concordance with tumor tissue and plasma cfDNA while the BRAF mutation analysis showed a 100% concordance [23]. Very recently, Sclafani et al. [142] analyzed the most frequently mutated KRAS hotspot mutation (i.e., G12D, G12V, and G13D) in rectal cancer tissue and corresponding plasma and did not observe much of a difference in the rate of KRAS mutation in plasma between patients with wild-type KRAS and KRAS-mutated tumors. Interestingly, by analyzing of plasma ctDNA the presence of additional KRAS mutations that were not detected on tissue were identified. Bettegowda et al. [21] detected that the sensitivity of ctDNA for the detection of clinically relevant KRAS gene mutations was 87% for stage IV CRC patients while in stage I CRC patients, a decrease to 47% was observed. Yang et al. [152] monitored that the ctDNA concentration increased with tumor size and cancer stage, where Stage IV patients evinced higher ctDNA levels than Stage I patients. Surprisingly, the Stage IV CRC patients showed only slightly over one mutated gene per patient than those with CRC Stage I.

Epigenetic analysis of cfDNA might contribute to the identification of gene hypermethylation [86] or the type of cell that leads to a rise of cfDNA fragments [163], and thus provide information about the tumor microenvironment which usually lacks somatic mutations. The methylation status of cfDNA was studied less intensively [103,122].

Recently, Li et al. aimed to prove the possibility of detecting copy-number variation (CNV) in the serum and in plasma of CRC patients, individuals with polyps and healthy volunteers. The analysis showed more prominent copy number changes in plasma than in serum. From 80 patients with CRC, CNV was detected in 39 of them and the majority of them were patients with an advanced stage. The most common changes included whole chromosomes gains on chromosomes 2, 7, 13, and 20; partial gains at chromosomes 8, 12, 13, and 20; and partial losses at chromosomes 1, 3, 4, 8, 17, 18, and 22 [135].

2.2.2. Cell-Free DNA as a Predictive Biomarker in Colorectal Cancer

Proper information on the clinical/pathological staging and the possibility of identifying cancer patients with a high likelihood of recurrence, or risk of clinical toxicity, are vital for the development of more efficient/less toxic treatment strategies.
The treatment of CRC has advanced over the past several years with the introduction of several active agents. Determining which patients to treat with chemotherapy and choosing the optimal treatment regime would allow oncologists to maximize the benefit of chemotherapy. Several prognostic and predictive markers have been identified and include oncogenes, tumor suppressor genes, genes involved in angiogenic and apoptotic pathways and cell proliferation, and those encoding targets of chemotherapy.

Zitt et al. [101] evaluated whether the concentration of cfDNA has the potential to serve as a marker for therapy monitoring during the treatment course of locally advanced rectal cancer patients. As the standard treatment for locally advanced RC, preoperative chemoradiotherapy (CRT) is considered. The changes in cfDNA concentration might be also observed before, after, and at the end of the preoperative CRT. The authors divided their population into responders (T0–T2 stage) and non-responders (T3–T4 stage). Both groups evinced a similar median plasma cfDNA concentration before and after the end of CRT. At the end of the treatment, the responders exhibited a decrease in cfDNA, whereas, in non-responders, the cfDNA increased. This is in agreement with other cancer research stories, in which the authors demonstrated that the response is associated with a decrease in the plasma cfDNA level, whereas no change or even an increase in the amount of cfDNA was seen in patients who did not respond to the therapy [25,164,165].

In a recent report on four locally advanced rectal cancer patients whose ctDNA was tracked in serial blood samples by using two patient-specific chromosomal rearrangements, Carpinetti et al. [121] showed an overall lack of correlation between the normalization of ctDNA and the amount of the residual disease in the surgical specimens after neoadjuvant CRT. However, changes in ctDNA levels after surgery appeared to predict the tumor recurrence.

In 2002, Diaz et al. [105] studied whether mutant KRAS ctDNA could be detected in the serum of CRC patients receiving monotherapy with panitumumab, a therapeutic anti-EGFR antibody. Authors found that 9 out of 24 patients whose tumors were initially without KRAS mutations developed detectable mutations in the KRAS gene in their sera. This suggests that KRAS mutations act as a mediator in acquired resistance to EGFR blockade and that these mutations can be detected in a non-invasive manner. Mohan et al. [115] reported the acquirement of resistance to anti-EGFR therapy with developed KRAS mutations and that resistant clones in the bloodstream had been detectable several months before the progression clinically manifested. A similar was we observed in the studies of Valtorta et al. [166], Misale et al. [167], and Bardelli et al. [168]. Another study focused on the abundance of mutant KRAS/BRAF alleles in the plasma of metastatic CRC patients at the baseline and before each cycle of third-line treatment with cetuximab and irinotecan [117]. cfDNA and KRAS levels decreased from the baseline to cycle 3 and increased at the time of progression. The decrease was larger in responding patients than in non-responding patients. The same group also investigated the total cfDNA in CRC patients during treatment with second-line chemotherapy and cfDNA in healthy controls [118]. cfDNA levels were significantly higher in CRC compared to controls. Patients with high cfDNA levels had a shorter survival after irinotecan-based therapy compared to those with lower levels.

The cfDNA integrity number (i.e., a ratio between long and short DNA fragments) turned out as an independent marker on CRT. The cfDNA integrity index was lower in responders compared to non-responders after CRT (as defined by the degree of tumor regression according to the Mandard score) only after completion of fluoropyrimidine-based CRT and, generally, patients with CRC had a 10-times higher cfDNA integrity number than in healthy subjects [104]. The DNA integrity index was observed to be the independent predictive factor of response to neoadjuvant treatment in multivariate analysis. Sun et al. [113,169] confirmed the potential of the DNA integrity index as they observed an association between this parameter (both at the baseline and after neoadjuvant treatment) and tumor regression grading according to the Dworak’s score in rectal cancer patients who received an oxaliplatin-based CRT for cT3-4 and/or N+ rectal tumors. The authors also observed that the plasma level of the KRAS mutation (codon 12) decreased with CRT in all patients with no difference between
responders and non-responders and that the higher MGMT promoter methylation status at baseline DNA was associated with a better tumor response.

2.2.3. Cell-Free DNA as a Prognostic Biomarker in Colorectal Cancer

The prognosis of patients with CRC is highly impacted by various factors at the time of diagnosis (tumor localization, quality of surgical procedures, gender, age, and the patient’s overall performance status).

Recently, the prognostic and/or predictive values of ctDNA in CRC have attracted the most intense interest. It is hypothesized that the mutational pattern observed in ctDNA might aid to stratify patients’ group into molecular subtypes with different prognoses, for example in diffuse large B lymphoma [169] or breast cancer [170]. Yi et al. [170] observed that the number of somatic mutations increased after therapy and the fractions of truck mutations were positively associated with targeted therapy. Studies uncovered that presence of ctDNA could be a reliable prognostic factor correlated with poorer outcome and support the idea of ctDNA as a non-invasive biomarker of minimal residual disease (Table 3.). Cassinotti et al. [110] and Frattini et al. [92] observed that after primary resection, the concentration of cfDNA significantly decreased, however, in patients with a relapse, the cfDNA levels dramatically increased, while, in “disease-free” patients, the cfDNA concentration still had a decreasing tendency. The preoperative measurement of cfDNA might contribute to the better estimation of prognosis and a post-operative measurement could represent a promising tool for the early detection of recurrence. A higher cfDNA concentration significantly correlated with a worse survival in other studies [100,126]. Diehl et al. [97] observed that CRC patients with detectable ctDNA after surgery generally relapsed within 1 year and considered ctDNA as a highly specific biomarker of tumor dynamics. This might indicate that ctDNA levels in plasma after surgery represent a reliable marker of residual disease. The utility of serial samplings proved the assumption of relapse prediction. Reinert et al. [123] observed a 2–15 (mean 10) months lead time on detection of metastatic recurrence compared to conventional follow-up. El Messaoudi et al. [129] postulated that all cfDNA parameters might be considered as prognostic markers: patients with higher levels of mutant ctDNA and higher mutation loads for the detected mutations in KRAS or BRAF genes evinced shorter OS. Similarly, Xu et al. [120] observed that patients with KRAS mutations evinced worse overall survival (OS) than KRAS wild-type patients. Higher ctDNA levels were associated with a higher risk of recurrence and a worse OS in CRC patients treated with surgery, chemotherapy, radiotherapy, or targeted therapy [87,112].

The detection of ctDNA in CRC patients with stage I, II, or III was associated with a shorter survival and disease recurrence. Lecomte et al. [86] observed a decreased 2-year survival in patients with cfDNA in plasma when compared to those without cfDNA in their plasma and suggested that patients with a high risk of recurrence can be distinguished by the analysis of tumor-derived ctDNA. Wang et al. [27] analyzed the presence of mutations in APC, KRAS, and TP53 genes in the serum of CRC patients and observed a significant correlation between the ctDNA and developed postoperative recurrence. Similarly, Tie et al. [130] in stage II CRC patients noticed that ctDNA detected postoperatively increased when not treated with chemotherapy. On the other hand, patients with negative ctDNA postoperatively were at a lower risk of radiological recurrence. Similarly, authors observed that ctDNA measurements seem to be a more sensitive marker of recurrence prediction than CEA levels. More than 80% of CRC patients were ctDNA positive at the time of recurrence, while only 41% of CRC patients evinced elevated CEA levels. Lately, Reinert et al. [123] and Carpinetti et al. [121] found that ctDNA monitoring in CRC patients may render an earlier cancer recurrence and therapy response in comparison with standard CEA measurements or radiological diagnosis. Similarly, Yang et al. [152] detected elevated CEA levels in 20% of Stage I CRC patients in comparison to 86% of Stage IV patients, while ctDNA levels above the limit 0.01 ng/µL were detected in 96% of all CRC patients (including Stage I). This hallmark points to the idea that ctDNA as
a supplement to traditional diagnostics with real-time tumor information and shows a much greater sensitivity than other tumor biomarkers.

The presence of methylation of HLTF and HPP1 genes was associated with a worse survival [94,107]. Lee et al. [111] analyzed the promoter methylation of the Septin 9 gene among CRC patients’ stage I-II and suggested that the methylation of the Septin 9 might be associated with lower disease-free survival. Recently, Herbst et al. [133] suggested that the detection of HPP1 methylation in cfDNA might be used as a prognostic marker and an early marker to identify patients who will likely benefit from a combination of chemotherapy and bevacizumab.

In summary, Fan et al. [171], in their comprehensive review, revealed that a ctDNA-positive status is associated with a worse prognosis. The authors performed a systematic review of data from published studies until December 2016.

2.3. Cell-Free DNA in Other Body Fluids

The liquid biopsy research is continuously developing. Many researchers are looking for new non-invasive approaches to detect and analyze ctDNA in other body fluids. Fujii et al. [172] monitored 56 patients with CRC KRAS mutations in their urine, plasma, and archival tumor tissue using mutation enrichment PCR coupled with NGS [172]. The concordance between ctDNA in urine and mutant KRAS in the tumor was 89% (sensitivity 80%, specificity 100%). In their study, patients had significantly fewer mutant cfDNA KRAS copies in urine during systemic therapy than at baseline or disease progression. Compared with no changes or increases in urine mutant cfDNA KRAS copies during therapy, decreases in these measures were associated with a longer median time to treatment failure. These preliminary findings suggest that ctDNA in urine may also reflect tumor dynamics and serve as a valid method of monitoring during treatment.

Several authors focused their study on the identification of stool biomarkers in CRC research. Ahlquist et al. [173] examined the possibility of cfDNA alterations in stools to discriminate between subjects with CRC and healthy individuals. APC, TP53, and KRAS gene mutations and BAT26 microsatellite instability were evaluated and the authors reported a 91% sensitivity and 93% specificity in detecting CRC. The same authors, Ahlquist et al. [174], used a next-generation stool DNA test based on a quantitative allele-specific real-time target and signal amplification assay with the aim of detecting early-stage CRC. The sensitivity of 85% for CRC and a specificity of 89% were obtained.

3. Conclusions

The aim of this review was to describe the different aspects related to cfDNA/ctDNA and evidence from studies supporting its potential use in CRC diagnoses and the improvement of therapy efficacy and prognosis. Most of the articles found that the presence of ctDNA in serum or plasma is associated with a worse survival possibility for patients with CRC.

The potential of the liquid biopsy in CRC research is attracting many scientists and has started to be a part of several clinical trials. However, before implementation of the liquid biopsy in clinical practice, it is necessary to set up standardized pre-analytical methodologies, including blood collection, processing, and storage and DNA extraction, quantification and, of course, validation in large prospective clinical studies [12]. Similarly, a better understanding of the origin and biology of cfDNA and ctDNA would help in the implementation of the results [175]. The impact of apoptosis and necrosis and the active release should be also explored. Our limited understanding of the release and clearance mechanisms of cfDNA hinders the interpretation of the current research [58].

Despite the ctDNA analysis having a great sensitivity and specificity compared with conventional diagnostic tools, taking a multi-marker approach may offer a more comprehensive insight into patient prognoses or therapy responses [23,59,129].

Since ctDNA provides real-time molecular information to track treatment response and relapse, it has a big potential to unravel drug-resistance mechanisms. With the advantage of next-generation
sequencing, multiple mutations in ctDNA could be identified. Precision medicine may change the clinical practice by adapting treatment choices based on an individual’s genetic background.

In the future, clinical trials on the ctDNA-based decision made in therapy selection would be definitive. Several trials of the clinical utility of ctDNA analyses for treatment monitoring are now being carrying out on NSCLC and breast cancer patients [176,177]. In summary, this emphasizes the fact that ctDNA based research is translating from exploratory research towards clinical trials in which ctDNA is acting as a decision-making tool.

These all support the further development of ctDNA as a biomarker for the detection of early diseases and minimal residual diseases following curative resection, relapse assessment, treatment response, and the development of chemoresistance. As a summary, we can conclude that the liquid biopsy should be considered as key towards the introduction of personalized medicine and patient benefits.

4. Methods

We searched the Ovid MEDLINE and PubMed databases using cell-free DNA, circulating tumor DNA, liquid biopsy and colorectal, colon and rectal cancer as keywords without any exclusion. We also searched bibliographies manually.

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Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| ARMs         | amplification refractory mutations system |
| BEAMing      | bead emulsion amplification and magnets |
| CAPP-Seq     | cancer personalized profiling by deep sequencing |
| cSMART       | circulating single molecule amplification and re-sequencing technology |
| ctDNA        | circulating cell-free DNA |
| COLD-PCR     | Co-amplification at lower denaturation temperature |
| CNV          | copy number variations |
| CRC          | colorectal cancer |
| CRT          | chemoradiotherapy |
| CTC          | circulating tumor cells |
| ctDNA        | circulating tumor DNA |
| ddPCR        | droplet digital PCR |
| DDR          | DNA-damage response |
| dPCR         | digital PCR |
| FISH         | fluorescent in situ hybridization |
| MSI          | microsatellite instability |
| MS-PCR       | methylation specific PCR |
| NGS          | next-generation sequencing |
| PNAs-LNA     | peptide nucleic acid-locked nucleic acid |
| qPCR         | quantitative PCR |
| RC           | rectal cancer |
| RT-PCR       | real-time PCR |
| Safe-Seq     | safe sequencing system |
| SERS         | surface-enhanced Raman spectroscopy |
| SNV          | single nucleotide variation |
| SOLID        | Sequencing by Oligonucleotide Ligation and Detection |
| SSCP         | single strand conformation PCR |
| TAM-Seq      | tagged-amplicon deep sequencing |
| UltraSEEK    | high-throughput multiplex ultrasensitive mutation detection |
| WGS          | whole genome sequencing |
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