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Quantifying circulating cell-free DNA in humans

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To our knowledge, this is the first comprehensive study on the influence of several pre-analytical and demographic parameters that could be a source of variability in the quantification of nuclear and mitochondrial circulating DNA (NcirDNA and McirDNA). We report data from a total of 222 subjects, 104 healthy individuals and 118 metastatic colorectal cancer (mCRC) patients. Approximately 50,000 and 3,000-fold more mitochondrial than nuclear genome copies were found in the plasma of healthy individuals and mCRC patients, respectively. In healthy individuals, NcirDNA concentration was statistically influenced by age (p = 0.009) and gender (p = 0.048). Multivariate analysis with logistic regression specified that age over 47 years-old was predictive to have higher NcirDNA concentration (OR = 2.41; p = 0.033). McirDNA concentration was independent of age and gender in healthy individuals. In mCRC patients, NcirDNA and McirDNA levels were independent of age, gender, delay between food intake and blood collection, and plasma aspect, either with univariate or multivariate analysis. Nonetheless, ad hoc study suggested that menopause and blood collection time might have tendency to influence cirDNA quantification. In addition, high significant statistical differences were found between mCRC patients and healthy individuals for NcirDNA (p < 0.0001), McirDNA (p < 0.0001) and McirDNA/NcirDNA ratio (p < 0.0001). NcirDNA and McirDNA levels do not vary in the same way with regards to cancer vs healthy status, pre-analytical and demographic factors.

Since Mandel and Metais discovered the presence of nucleic acids in serum in the 1940s1, different studies have reported elevated levels of circulating DNA (cirDNA) in the blood of patients suffering from various diseases2-6, especially cancer7-10. Despite scant early consideration, interest in the feasibility of cirDNA analysis has increased exponentially, over the last decade, among researchers working on a large range of disorders. CirDNA was first clinically implemented in prenatal diagnosis of sex-determination and pregnancy-associated disorders by assay-
affect its measurement\textsuperscript{16}. Human cellular aging is usually marked by senescence and cell death. Similarly, it often features a phenomenon, originally called “Inflamm-aging”, that induces a chronic or low-grade inflammatory state\textsuperscript{17–20}. Indeed, it was demonstrated that tissue damage and a pro-inflammatory environment increase release of cirDNA into the blood stream\textsuperscript{14}. Additionally, we may speculate that sex-based differences, such as genetic dissimilarity and steroid hormones levels, could cause differences in cirDNA concentration between men and women. Likewise, blood component concentrations may vary with the circadian clock and upon food intake and potentially influencing cirDNA concentration\textsuperscript{21}. Despite outstanding research in the field of cirDNA, relatively few studies have examined the effect of pre-analytical and demographic parameters as sources of intra- and inter-individual variability in cirDNA levels. There is currently no single operating procedure and there are relatively few clinical guidelines in the literature\textsuperscript{22–25}. This study aims first at defining a framework for cirDNA analysis, to harmonize NcirDNA and McirDNA quantification and to explore potential sources of variability that could cause interpretation errors. Previous studies already documented various pre-analytical limitations and specified conditions for cirDNA analysis, including specific collection tubes, and specific plasma isolation and extraction protocols, including storage condition variables and limits on the number of freeze-thaw cycles\textsuperscript{22–24,26}. Here, we study the influence of various pre-analytical (plasma aspect, delay between blood collection and last food intake) and demographic variables such age and gender in NcirDNA and McirDNA concentration determination, from plasma of 104 healthy donors and 118 mCRC patients (Fig. 1). In addition, we compared McirDNA and NcirDNA levels in healthy individuals and mCRC patients. Some others pre-analytical conditions such as blood collection tubes, blood collection time and blood stability were examined \textit{ad hoc} study on healthy volunteers.

\section*{Results}

\subsection*{Simultaneous quantification of NcirDNA and McirDNA.}

We used serial dilutions of genomic DNA and of mitochondrial plasmid DNA to validate a reproducible, linear and sensitive assay to quantify, both nuclear and mitochondrial, circulating DNA (NcirDNA and McirDNA). The NcirDNA assay is based upon the qPCR detection of a sequence of the \textit{KRAS} gene and it can detect one copy of nuclear \textit{KRAS} gene per 6 microliters of plasma (1 nuclear Genome Equivalent per 12 \(\mu\)L). The McirDNA assay can detect down to one copy of the mitochondrial \textit{MT-CO3} gene per 1.7 microliter of plasma (1 mitochondrial Genome Equivalent per 1.7 \(\mu\)L). Note, the targeted \textit{MT-CO3} gene sequence was selected as being not mutated in the mitochondrial genome of cancer patients. Copy number calculation is performed by using a specific equation that eliminates bias in the nuclear or mitochondrial DNA calibration curves to allow simultaneous calculation of their real relative proportions. For example, in the cohort of 104 healthy individuals, we found a median NcirDNA and McirDNA plasma concentration of \(1.64 \times 10^3\) and \(8.32 \times 10^7\) copies/mL, respectively, corresponding to 5.43 and 1.36 ng/mL of plasma.

\subsection*{Comparison between NcirDNA and McirDNA levels.}

We compared NcirDNA and McirDNA concentration, in healthy individuals and mCRC patients groups, expressed either in copy number/ml and ng/ml of plasma. A supplementary table summarizes all quantification data and comparative tests in detail (Supplementary Table S1). There was a highly statistical difference between NcirDNA and McirDNA concentration expressed in copy number/ml, in healthy individuals (Fig. 2A; \textit{Mann-Whitney U test, \(P\) value < 0.0001) and equally in mCRC patients (Fig. 2B; \textit{Mann-Whitney U test, \(P\) value < 0.0001). We next compared McirDNA and NcirDNA concentration expressed in ng/ml, in healthy individuals and mCRC patients groups. Here, also, we observed a considerable statistical difference between NcirDNA and McirDNA concentrations in healthy individuals (Fig. 2C; \textit{Mann-Whitney U test, \(P\) value < 0.0001) and mCRC patients (Fig. 2D; \textit{Mann-Whitney U test, \(P\) value < 0.0001). McirDNA concentration is significantly higher than NcirDNA concentration when measured in units of copy number/ml, and conversely, the NcirDNA concentration was significantly higher than the McirDNA concentration when measured in ng/ml, either in healthy individuals and mCRC patients.
Effect of age and gender on cirDNA concentrations in healthy individuals. **NcirDNA.** **Age** (n = 104): We dichotomized the healthy individuals cohort in two groups around the median age, which was 47 (Table 1). The median NcirDNA copy number in the <47 years-old group (n = 52) and in the ≥47 years-old group (n = 52) were 1.36 × 10^3 and 1.73 × 10^3 copies/ml, respectively. A statistical difference was found between young and older healthy individuals groups (Fig. 3A; Mann-Whitney U test, P value = 0.009).

**Gender** (n = 104): The median NcirDNA copy number in the healthy male group (n = 62) and in the group of healthy females (n = 42) were 1.69 × 10^3 and 1.48 × 10^3 copies/ml, respectively. This difference in the NcirDNA copy number between healthy males and females was statistically significant (Fig. 3B; Mann-Whitney U test, P value = 0.048).

**Multivariate analysis:** Logistic regression analysis including age and gender specified that age over 47 years-old was predictive to have higher NcirDNA concentration (Fig. 3C; OR = 2.41, P value = 0.033). Results of multivariate analysis and Odds ratios (OR) with 95% confidence intervals (CIs) are summarized in Supplementary Table S2A.

**McirDNA.** **Age** (n = 104): The median McirDNA copy number in the <47 (n = 52) and in the ≥47 year-old group (n = 52) were 7.77 × 10^7 and 8.40 × 10^7 copies/ml, respectively. There was no statistical difference in the McirDNA copy number between these groups (Fig. 3D; Mann-Whitney U test, P value = 0.489).

**Gender** (n = 104): The median McirDNA copy number in the healthy male group (n = 62) and in the group of healthy women (n = 42) were 8.03 × 10^7 and 9.39 × 10^7 copies/ml, respectively. No statistical difference in McirDNA copy number was observed between healthy males and females (Fig. 3E; Mann-Whitney U test, P value = 0.485).

**Multivariate analysis:** Logistic regression analysis including age and gender confirmed no statistically significant difference between studied groups (Fig. 3F). Results of multivariate analysis and Odds ratios (OR) with 95% confidence intervals (CIs) are summarized in Supplementary Table S2B.
Effect of various parameters on cirDNA concentrations in mCRC patients. **NcirDNA. Plasma aspect (n = 106):** We first compared two groups, 27 abnormal plasmas (icteric and/or opaque plasmas) and 79 normal plasmas. There was no statistical difference in NcirDNA levels between abnormal plasmas and normal plasmas groups (Fig. 4A; Mann-Whitney U test; P value = 0.266). However, the median NcirDNA amount determined in abnormal plasmas group was slightly lower than in normal plasmas (3.62 × 10^3 vs 5.22 × 10^3 copies/ml).

**Delay between blood collection time and last food intake (n = 89):** We then compared two groups, “1 h < delay < 5 h” (n = 69) and “other delays” (n = 20). There was no statistical difference in NcirDNA levels between “1 h < delay < 5 h” and “other delays” (Fig. 4B; Mann-Whitney U test; P value = 0.220). Nonetheless, the “1 h < delay < 5 h” group showed lower median NcirDNA amount than the group “other delays” (3.68 × 10^3 vs 5.72 × 10^3 copies/ml).

**Age (n = 118):** We also dichotomized the mCRC cohort into two groups around the median age of 65 (Table 1). The median NcirDNA copy number in the <65 year-old group (n = 52) and in the ≥65 year-old group (n = 66) were 4.73 × 10^3 and 4.94 × 10^3 copies/ml, respectively. No statistical difference was found (Fig. 4C; Mann-Whitney U test, P value = 0.757). A comparative study using the same cut-off for both healthy and mCRC cohorts as the median age of all individuals tested here (N = 222, median age = 56 years) confirmed the statistical difference between young (N = 79) and older (N = 25) healthy individuals groups (Mann-Whitney U test, P value = 0.0026), and also the no statistical difference between young (N = 25) and older (N = 93) mCRC groups (Mann-Whitney U test, P value = 0.913) (Supplementary Fig. S1A,B).

**Gender (n = 118):** The median NcirDNA copy number in the mCRC males (n = 68) and in and females (n = 50) mCRC patients were 4.65 × 10^3 and 5.40 × 10^3 copies/ml, respectively. No statistical difference was found (Fig. 4D; Mann-Whitney U test, P value = 0.971).

**Multivariate analysis:** Logistic regression analysis including all the parameters (plasma aspect, delay, age and gender) confirmed no statistically significant results (Fig. 4E). Results of multivariate analysis and Odds ratios (OR) with 95% confidence intervals (CIs) are summarized in Supplementary Table S2C.

**McirDNA. Plasma aspect (n = 46):** We next compared the abnormal plasmas (n = 11) and normal plasmas (n = 35) groups. There was no statistical difference in NcirDNA levels between abnormal and normal plasmas groups (Fig. 5A; Mann-Whitney U test; P value = 0.263). However, the median McirDNA amount determined in abnormal plasmas was slightly lower than in normal plasmas (6.64 × 10^3 vs 1.19 × 10^4 copies/ml).

**Delay between day-time of blood draw and the last food intake (n = 34):** There was no statistical difference in McirDNA copy number between the “1 h < delay < 5 h” group (n = 25) and the “other delay” group (n = 9) (Fig. 5B; Mann-Whitney U test; P value = 0.509). The “1 h < delay < 5 h” group showed higher median NcirDNA amount than the group “other delay” (1.68 × 10^3 vs 1.19 × 10^4 copies/ml).

**Age (n = 50):** The median McirDNA copy number in the <65 year-old (n = 23) and ≥65 year-old (n = 27) groups were 1.12 × 10^4 and 1.84 × 10^4 copies/ml, respectively. There was no statistical difference in the McirDNA copy number between the groups (Fig. 5C; Mann-Whitney U test, P value = 0.240).

**Gender (n = 50):** The median McirDNA copy number in the mCRC males group (n = 25) and in the group of mCRC women (n = 25) were 1.19 × 10^4 and 1.68 × 10^4 copies/ml, respectively. We did not found any statistical difference in the McirDNA copy number between mCRC males and females (Fig. 5D; Mann-Whitney U test, P value = 0.261).

**Multivariate analysis:** Logistic regression analysis including all the parameters (plasma aspect, delay, age and gender) confirmed no statistically significant results (Fig. 5E). Results of multivariate analysis and Odds ratios (OR) with 95% confidence intervals (CIs) are summarized in Supplementary Table S2D.

| Patient's characteristics | Healthy individuals (N = 104) | mCRC patients (N = 118) |
|---------------------------|------------------------------|------------------------|
| Age (years)               | Mean 45                      | Mean 65                |
| Median                    | 47                           | 65                     |
| (min-max)                 | (18–69)                      | (22–91)                |
| Gender                    |                              |                        |
| Males                     | 62                           | 68                     |
| Females                   | 42                           | 50                     |
| TOTAL                     | 99                           | 118                    |
| Mean age                  | 45                           | 65                     |
| Median age                | 47                           | 65                     |
| (min-max)                 | (19–69)                      | (34–88)                |
| Females (N)               |                              |                        |
| Mean age                  | 44                           | 65                     |
| Median age                | 45                           | 67                     |
| (min-max)                 | (18–63)                      | (22–91)                |

Table 1. Characteristics of healthy individuals (N = 104) and mCRC patients (N = 118).
Comparing cirDNA levels between mCRC patients and healthy individuals. 

**NcirDNA.** We compared the median NcirDNA amount between healthy individuals (n = 104) and mCRC patients (n = 118). The median NcirDNA concentration in healthy individuals and mCRC patients was $1.64 \times 10^3$ and $4.73 \times 10^3$ copies/ml, respectively, revealing a significant difference between healthy individuals and mCRC patients (Fig. 6A; Mann-Whitney U test, P value < 0.0001).
Next we compared the median McirDNA amount between healthy individuals (n = 104) and mCRC group (n = 50). The median McirDNA concentration in healthy individuals and mCRC patients was statistically different at $8.32 \times 10^7$ and $1.44 \times 10^7$ copies/ml, respectively (Fig. 6B; Mann-Whitney U test, $P$ value $<0.0001$).

**McirDNA/NcirDNA ratio.** Finally we compared the median McirDNA/NcirDNA ratio between healthy individuals (n = 104) and mCRC patients (n = 50). The median McirDNA/NcirDNA ratio in healthy individuals

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**Figure 4.** Influence of various factors on NcirDNA concentration in mCRC patients. Boxplot analysis of NcirDNA concentration extracted from mCRC patients (N = 118), with regards to plasma aspect (A); delay between blood collection and last food intake (B); age (C) and gender (D). (E) Multivariate analysis representation. Boxplot represent median with min to max of values. Mann-Whitney U test was performed for univariate analysis and logistic regression was performed for multivariate analysis. Odds ratio (OR) with 95% confidence intervals (CIs) are presented. A probability of $\leq 0.05$ was considered to be statistically significant; $^*p \leq 0.05$, $^{**}p \leq 0.01$, $^{***}p \leq 0.001$, $^{****}p \leq 0.0001$. 

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**McirDNA.**
and mCRC patients were statistically different at $5.41 \times 10^4$ and $2.70 \times 10^3$ copies/ml, respectively (Fig. 6C; Mann-Whitney U test, $P$ value $< 0.0001$). The median McirDNA/NcirDNA ratio was 20-fold higher in the healthy individuals group than in the mCRC patients group. We did the same comparison using McirDNA and NcirDNA concentrations expressed in ng/ml and we obtained the same results (Supplementary Fig. S2, Mann-Whitney U test, $P$ value $< 0.0001$).
**CirDNA stability in whole blood samples of healthy volunteers.** NcirDNA concentration in blood collected in EDTA tubes increased by more than two-fold and three-fold respectively, at days 2 and 5 after blood collection (Supplementary Fig. S3A). In contrast, there was no difference between days 0, 2, 5 and 7 following
blood collection in Cell-Free DNA BCT® STRECK (BCT) tubes (Supplementary Fig. S3B). Note, median values at day 0 are similar in plasma from blood collected in EDTA or BCT tubes. In addition, the McirDNA concentration in blood collected in EDTA tubes approximately increased more than 12-fold and 27-fold respectively, at days 2 and 5 after blood collection (Supplementary Fig. S3C).

**Effect of the blood collection time on cirDNA levels in healthy volunteers.** NcirDNA. In EDTA tubes at 9.00 AM (fasted state), 12.00 PM (2 hours after breakfast), 3.00 PM (2 hours after lunch) and 6.00 PM, the median NcirDNA copy numbers were 1500, 480, 715 and 790 copies/ml respectively. An additional figure shows this in more detail (Supplementary Fig. S4A,E). Taking the four healthy individuals together, the median NcirDNA concentration strongly changed from the fasted state to three hours after breakfast. Median NcirDNA concentration was slightly higher at the earliest collection time in BCT tubes, over the same time-course of the day, at 874, 562, 728 and 313 copies/ml, respectively. An additional figure shows this in more detail (Supplementary Fig. S4B,F). Altogether our data from both collection methods showed that NcirDNA content in healthy individuals declined from 9.00 AM to 12.00 PM then plateaued up to the 6.00 PM collection time. An additional figure shows this in more detail (Supplementary Fig. S4C,D,G,H).

McirDNA. In EDTA tubes from healthy individuals, the median McirDNA copy numbers were unchanged over the time course at 1.05, 1.25, 1.17 and 1.26 million copies/ml. An additional figure shows this in more detail (Supplementary Fig. S5A,B). No clear difference was found when comparing McirDNA amounts collected at the various collection time points. Note, plasma appearance become opaque at 6.00 PM. An additional picture illustrates this observation (Supplementary Fig. S6).

**Discussion**

Here, we present a comprehensive study on the quantification of nuclear and mitochondrial cirDNA in the plasma of healthy individuals and a homogenous cohort of mCRC patients. To our knowledge, this is the first to address altogether the influence of various preanalytical, analytical and demographical factors on NcirDNA and McirDNA levels in a large set of individuals (n > 200). Solid tumor mass is composed of a variety of cells, mostly consisting of malignant/cancer, stromal, endothelial and immunological cells. In order to avoid any confusion in nomenclature, we define as ‘tumor cells’ all the cells composing the tumor mass. The concentration values presented in this study correspond to total cell-free DNA in plasma, either of nuclear or mitochondrial origin. Our observations are summarized in Table 2 and Supplementary Table S3.

**Total cirDNA levels.** This study design relies on the analytical performance of the assay. We used here a qPCR-based method with an unmatched combination, of reproducibility, sensitivity and specificity for quantifying plasma cirDNA (see Methods section). Furthermore, our assay benefits from a clinically-validated optimal pre-analytic process that we previously set up for plasma preparation27, and adapted from Chiu et al., DNA extraction and sample handling27. The accuracy of the cirDNA concentration measurement in this study is supported by two assessments: (i) total cirDNA concentration by targeting a KRAS sequence was routinely controlled by quantifying a BRAF sequence. In addition, this control quality enable to detect and exclude sample with loss of heterozygosity (LOH) or gene amplification which have been reported in CRC patients28. Moreover, since KRAS amplification is an infrequent event in CRC (0.67%)29, this level will not modify our observations or the values described in our manuscript; and (ii) the study of cirDNA measurement under Poisson law distribution revealed single copy detection of nuclear cirDNA.

Respective NcirDNA and McirDNA proportions inverted when the plasma cirDNA concentrations were calculated according to copy number or mass (Fig. 2A,C). For example, data obtained from 104 healthy individuals revealed that median McirDNA plasma concentration was approximately 50,000-fold higher than median NcirDNA plasma concentration in terms of copy number/ml (99.998% McirDNA and 0.002% NcirDNA), corresponding to approximately 4-fold lower in terms of ng/ml (25.0% McirDNA and 75.0% NcirDNA). This highlights the much lower size of the mitochondrial genome (16569 bp) than the nuclear genome (about 3 × 10⁹ bp), and also the high number of mitochondrial genome copies within a cell. Each human cell, depending on its type, contains a number of mitochondria ranging from 500 to 2,000 and each mitochondria holds between 2 and 10 mitochondrial DNA molecules. Therefore, each cell may contain approximately 1,000 to 20,000 mitochondrial DNA molecules and only one nuclear genome equivalent, which is consistent with our results. Our method of calculation, described here, appears as the most rigorous means to simultaneously quantify mitochondrial and nuclear DNA concentrations. A higher coefficient of variation was routinely observed for mitochondrial than for nuclear DNA, despite the higher analytical signal for McirDNA copy number. The median concentration levels we found here appear to be similar to the average values observed in several studies27,28. In addition, performance of our assay support the accuracy of the values presented throughout this study such as 5.43 and 1.36 ng/mL in healthy subject plasma of N- and McirDNA, respectively. Considering the significant percentage of McirDNA (i.e. 25% in healthy subjects) among total cirDNA mass existing in plasma, we propose to always specify the origin of cirDNA when examining total circulating WT DNA.

**Influence of age and gender on NcirDNA levels in healthy individual plasma.** Most studies found no relationship between NcirDNA levels and any demographic parameters, such as age and gender, in healthy individuals30–32. A few studies showed opposite observations31–33. Here, we observed an influence of age on NcirDNA plasma concentration in the healthy individual cohort as a whole (p = 0.009). Our data also revealed a statistical difference in NcirDNA amount between healthy males and females (p = 0.048). We performed logistic regression for multivariate analysis including age and gender and showed that age ≥ 47 was predictive of a high NcirDNA concentration (OR = 2.41, p = 0.033). This observation is in accordance with a study by Jylhävä et al.
that consisted of 12 nonagenarian women (age > 90 years) and 11 healthy control female (22 < age < 37 years) that showed a higher concentration of NcirDNA in nonagenarians than in control women38. The authors explained this increase in the amount of cirDNA with age, as the accentuation of a senescence phenomenon and cell death, caused by an inflammation associated with age and even by decreased clearance and phagocytic capacity. Zhong et al. showed a significant increase in total plasma cirDNA concentration in women over 60, compared to younger women, which appears to be consistent with Jylhävä’s study. We may speculate that menopause could be an explanation of the statistically higher NcirDNA concentration in healthy males as compared to healthy females (p = 0.048), whereas no difference between mCRC males and females was observed (p = 0.971). This speculation is based on two observations: (i) Median menopausal age in the European population is 51 years-old; 10–15% of women go into menopause before the age of 45, and globally 90–95% reach menopause by 55 years of age. By applying these categories of age to our women cohorts, we observed no statistical difference and no tendency between mCRC women < 45, 45–54 and ≥ 55 year-old (Kruskal-Wallis rank test, P value = 0.592) while we showed a statistical difference between healthy women < 45, 45–54 and ≥ 55 year-old, with a concentration gradient: 1275, 1440 and 2610 median copy number, respectively (Kruskal-Wallis rank test, P value = 0.026); and (ii) CRC females are at a high risk of chemotherapy-induced menopause or menstrual disorders like amenorrhea or a decrease of steroid hormone levels36,37, and metachronous mCRC patients may attain a menopausal state earlier. An additional figure shows these results in more detail (Supplementary Fig. S7).

**Table 2. Summary of the observations made on the influence of demographical factors.**

Influence of various parameters on NcirDNA levels in mCRC patients plasma. Despite the large number of studies that aimed at determining if age and gender might influence NcirDNA levels, no clear results have been demonstrated30–32.38. There are discrepancies in the literature with regards to the influence of age and gender in patient populations suffering from various types of cancers. These discrepancies could result from use of serum, or pre-analytical or analytical factors. As an example, a study by Hohaus et al. showed that patients with Hodgkin and non-Hodgkin’s lymphoma over 60 year-old (n = 142) had higher levels of cirDNA in plasma than younger patients (p = 0.018)38. Conversely, our data showed no statistical influence of age (p = 0.757) and gender (p = 0.071) on NcirDNA concentration in mCRC patients and these results was confirmed with a multivariate analysis using logistic regression. We also demonstrated no influence of pre-analytical factors like plasma aspect (p = 0.266) or delay between last food intake and blood collection (p = 0.220). On the other hand, our results also confirm those previously published by our laboratory25 and by many teams39,40, namely that NcirDNA concentration is significantly higher cancer patients than in healthy individuals (p < 0.0001), whether male or female. For more than a decade it was suggested that total NcirDNA could be a cancer biomarker55,61,42. However, previous attempts to apply total cirDNA quantity as a screening test for cancer lacked a strong statistical demonstration55. High standards, with regard to pre-analytical factors and quantification, could lead to its use as one marker, among other, for tumor burden.

**McirDNA levels in plasma.** Relatively few reports have quantitatively analyzed McirDNA and there are discrepancies among them. In one study there was no significant difference between young and aged healthy subjects35. For Pinti et al. however, McirDNA concentration would increase with age48. In this study, McirDNA content was analyzed in 831 plasma samples from subjects with different healthy status, aged from 1 to 104 years; McirDNA content significantly increased after fifty years-of-age and it peaked in nonagenarians. Elevated McirDNA levels might
help maintain the low-grade chronic inflammation that is common in elderly individuals. With regards to McirDNA level in cancer patients, Mengel-From et al. measured McirDNA copy number in blood cells from 1,067 subjects aged 18 to 93 and conversely, observed a tendency for lower mitochondrial DNA copy number with advanced age44. These findings are consistent with other studies, performed on different types of tissue, such as skeletal muscle and pancreatic islets45,46. This age-related tissue-specific depletion of cellular mitochondrial DNA could lead to a proportional reduction of McirDNA copy-number in plasma. Inversely, there was no association between serum McirDNA levels and demographic parameters (age/gender) in urological malignancies47,48. Likewise, in breast cancer, there was no significant difference in McirDNA content in blood samples of stage I patients with respect to their age49. Our data revealed no influence of age and gender on McirDNA concentration, either in plasma of healthy individuals or mCRC patients. These results were confirmed in multivariate analysis. We also reported no significant influence of plasma aspect and delay between last food intake and blood collection (p = 0.509). Nonetheless, our data revealed a significant higher McirDNA than NcirDNA concentration in plasma, whether for mCRC patients, healthy individuals, male or female, and regardless of the age of the subject. Note, while median NcirDNA concentration is much higher in mCRC patients than in healthy individuals, median McirDNA concentration is conversely lower in mCRC patients, revealing a proportionally lower McirDNA release from cancer cells. This might be explained by the fact that cancer cells, in comparison to healthy cells, may have, among other differences, fewer mitochondria per cell and less DNA within their mitochondria44,50. However, this is still controversial and explanation of our striking observation is under active investigation in our team. Nevertheless, we may speculate that the McirDNA/NcirDNA ratio might have some power in discriminating healthy individuals from cancer patients. McirDNA/NcirDNA ratio is undergoing clinical validation as potential biomarker for tumor burden or diagnosis in a large study involving broader scope of cancer patients with various malignancies and stages. In light of the high copy number of McirDNA and its tendency to be mutated in cancer51, our observations confirm the gradual acceptance of McirDNA as a new potent diagnostic and prognostic biomarker for many solid tumors52.

Blood stability for cirDNA plasma assessment. As previously reported23,53, NcirDNA concentration determined from blood collected in EDTA tubes increased with time highlighting release of genomic DNA resulting from blood cell lysis when stored at room temperature or +4 °C and consequently to contamination of cell-derived DNA. Note, whole blood stored in EDTA tubes at +4°C showed no change in cirDNA concentration for up to one day suggesting their potential use within this time period (data not shown)23,53. We propose routine clinical analyses use plasma stored in EDTA tubes for up to 6 hours, given the uncertainty of maintaining the temperature of samples in the course of blood processing, as we earlier described35. Conversely, BCT tubes appeared to conserve blood cell integrity, since no DNA concentration increase was observed up to 7 days following blood collection23,53. Thus, BCT tubes maintain the true cirDNA concentration and are good tools to conserve/stabilize blood for optimal quantification of NcirDNA for up to 7 days following collection. Cell-preserving tubes greatly allows postal shipment of whole-blood within this time period and it allows interventional analysis as well as enabling clinical centers that lack lab facilities to immediately prepare plasma. While being cheaper by themselves, use of EDTA tubes necessitates plasma preparation within a short time frame and immediate subsequent storage under frozen conditions until analysis requiring costly shipment when plasma originate from a single patient. We first reported that McirDNA concentration determined from blood collected in EDTA tubes strongly increased with time. We may assume that this results as well blood cell lysis and blood cell-derived mitochondrial DNA contamination.

Effect of blood collection time on cirDNA plasma concentration. There is currently no indication in the literature on the optimal time for blood collection when analyzing cirDNA. Our data seem to indicate that NcirDNA median levels are 2- to 3-fold higher at 9.00 AM, which is the earliest time-point examined, compared to later blood-collection time-points (12.00, 3.00 and 6.00 PM) when the NcirDNA level stabilizes. Decrease from 9.00 AM to 12.00 PM might be explained by the postprandial effect of the breakfast being taken at 10.00 AM. This hypothesis is supported by several observations. First, we reported that NcirDNA plasma levels in blood collected between one and five hours after food intake were lower than in blood collected on patients under fasting conditions. It was previously showed that plasma triglyceride increased one hour after intake of a test meal and baseline values were restored back to initial values after 5 hours56. Second, NcirDNA concentration was lower in opaque than in non-opaque plasmas. Multivariate analysis including age, gender, plasma aspect and delay between food intake and blood collection revealed no statistical influence but abnormal plasma aspect showed a clear tendency to have lower NcirDNA concentration (OR = 0.399, p = 0.089). These observations are all consistent with postprandial effects. Food intake with high lipid content may result in hyperlipidemia which can be characterized by opaque plasma and high triglyceride concentrations. However, despite the large examined cohort data, we cannot state that postprandial is the explanation since no statistical difference was found. This may be due to various factors: (i) blood triglyceride levels largely depend on fat distribution and body weight, lifestyle choices, and also genetic factors57; (ii) there were considerable within- and between-subject variations in non-fasting plasma triglycerides46; and (iii) the subjects had a chronic illness, mCRC. We speculate that the postprandial effect could occur because the presence of lipids or proteins may interfere with DNA extraction yield from plasma. We cannot exclude the possibility that NcirDNA levels depend on circadian clocks and metabolism, resulting in more elevated concentration in the morning. Moreover, we cannot exclude the possibility that other metabolic changes during fasting/feeding alter cirDNA yield. Nevertheless, our data suggest that fasting blood samples should be included when studying or clinically examining cirDNA to improve its diagnostic performance, especially when low mutation frequency in cancer patients or prenatal testing is considered. Note, in addition to opaque plasma, we remarked on various occasions that, icteric plasma had aberrant cirDNA concentration when qPCR was the analytical method. Therefore, we propose observation of icteric, hemolytic and opaque plasma as criteria of blood sample exclusion.
Limitations of the study. While the study was carried out with statistically sufficient number of subjects to support the observations, the ad hoc study is limited by the low number of tested individuals since ad hoc study experiments are cumbersome and time-consuming. Thus, blood stability and blood collection time was only carried out on a few number of healthy volunteers (n = 5) and not on mCRC patients. This not allowed us to provide statistical analysis while the results showed clear tendencies. A specific study should be performed to statistically confirm these results on a larger cohort of healthy volunteers as well as mCRC patients. Although we routinely experienced that abnormal plasmas resulted in lowering cirDNA concentration values, we cannot fully discriminate the implication of postprandial effects like triglycerides serum level, to the possible involvement of biological changes due to the circadian rhythm. In order to definitively address this issue, it would be interesting to compare NcirDNA levels at 9.00 AM and 12.00 AM, with and without breakfast, in order to determine the impact of food intake and circadian rhythm, respectively. In addition, the influence of the menopause on cirDNA concentration with regard to our observations of gender, age and pathological status is only speculative. A specific study on the difference of NcirDNA concentration between postmenopausal and premenopausal healthy females, as well as the difference between premenstrual and postmenstrual young women, should be performed to definitively address this issue. Conclusions drawn here in respect to cancer plasma samples should be restricted to mCRC patients and extension to other malignancies or even to localized disease is speculative.

In conclusion, the levels of mitochondrial and nuclear circulating DNA differently vary with regards to pre-analytical and demographic factors. Those variables should be taken into consideration when evaluating cirDNA analysis in clinical setting and perhaps in the future clinical practice when cirDNA quantification is directly or indirectly used as a biomarker. In addition, our study highlights the potential for combining the analysis of NcirDNA and McirDNA since examining their respective levels may have diagnostic value.

Methods

Patients. Blood samples collected from 104 healthy donors were provided by the Etablissement Français du Sang (E.F.S), the blood transfusion center of Montpellier (Convention EFS-PM N° 21PLER2015-0013). Blood samples collected by the E.F.S were analyzed (virology, serology, immunology, blood numeration). If an anomaly is detected, the sample is ruled out and the donor is warned then by mail. NcirDNA concentration and McirDNA copy number were determined for all healthy individuals. Data on age and gender were collected for all healthy individuals. mCRC patients data are taken from a study comparing the detection of KRAS exon 2 and BRAF V600E mutations by circulating DNA (cirDNA) analysis to conventional detection by tumor tissue analysis39. This study (KPLEX2) was performed and presented under the STARD criteria. 140 patients have been included in 11 clinical centers in France, over a period of 12 months. Eligible patients were male or female, aged ≥18 years, with a proven histologically mCRC, a measurable disease as defined by response evaluation criteria in solid tumors (RECIST v1.1) and untreated by radiotherapy or chemotherapy in the last 4 weeks before inclusion. There is no possibility that cirDNA from mCRC patients can be affected by therapy since eligible patients were untreated by radiotherapy or chemotherapy in the last 4 weeks before blood collection (inclusion criteria). Written consent was obtained from the part of all patients. Inclusion criteria were described previously39. 19 were excluded from analysis for no respect of the inclusion criteria (due to various inclusion criteria) and 3 were excluded due to lack of data collected on age or gender. NcirDNA analysis was performed on 118 patients and McirDNA analysis on 50/118 mCRC patients. While study on age and gender effect were carried out on all the included mCRC patients for NcirDNA (N = 118), cohort patient number varied when studying plasma aspect and delay for NcirDNA, or age and gender for McirDNA because of two main reasons: non-reported information for delay and plasma aspect for to NcirDNA and insufficient plasma volume needed to carry out supplementary analysis for McirDNA. All data on age and gender were collected. Plasma aspect (normal, abnormal: opaque or/and icteric) was noted for 106/118 mCRC patients and delay between time of blood draw and the last food intake was informed for 89/118 mCRC patients.

Ethics approval and consent to participate. Blood samples collected from 104 healthy donors were provided by the Etablissement Français du Sang (E.F.S), the blood transfusion center of Montpellier (Convention EFS-PM N° 21PLER2015-0013). Plasma samples from mCRC patients were obtained from the Kpex2 study registration number EUDRACT 2016-001490-33 with ethic committee approval (“Comité de Protection des Personnes”, Nimes, France). All methods were performed in accordance with the relevant guidelines and regulations. The study obtained informed consent from all participants for the study.

Blood stability. 7 samples for each healthy donor were collected at 9.00 a.m. on Day 0 (fasted state): 3 with EDTA K2 tubes and 4 with Cell-Free DNA BCT® STRECK tubes. Each tube was processed as we early described, at day 0, 2 and 5 for EDTA tubes and day 0, 2, 5 and 7 for BCT tubes. EDTA tubes were stored at +4 °C and BCT tubes were conserved at room temperature before isolation. Healthy individual N°1 (HI1) and N°2 (HI2) are 57 year-old and 29 year-old men, respectively, both with no known disease. Healthy individual N°3 (HI3), N°4 (HI4) and N°5 (HI5) are 25 year-old, 24 year-old and 28 year-old women, respectively, both with no known diseases. NcirDNA analysis for HI1 was not performed due to clotting in BCT tubes.

Blood collection time. 4 blood collection times were defined 9.00 a.m. (fasted state), 12.00 p.m. (2 hours after breakfast), 3.00 p.m. (2 hours after lunch) and 6.00 p.m. (4 hours after lunch). One EDTA K2 and one Cell-Free DNA BCT® STRECK tube per day-time were collected for each donor. Healthy individual N°6 (HI6) and N°7 (HI7) are 29 year-old and 27 year-old men, respectively, both with no known diseases. Healthy individual N°8 (HI8) and N°9 (HI9) are 28 year-old and 30 year-old women respectively, both with no known diseases. Each donors took the same meal during breakfast: one butter croissant, one chocolate croissant and one coffee with sugar; and for the lunch: a dish of tomato rice with sausage, bread and one coffee with sugar.
Sample characteristics and preparation. Samples were collected and treated in accordance with a pre-analytical guideline previously established by our group. In summary, blood was collected in EDTA K3 tubes and plasma was isolated within 2 hours. The isolation technique consist of a double centrifugation. Initially, tubes were centrifuged for 10 minutes at 4°C and 1,200 g in a Heraeus Multifuge LR centrifuge. The supernatant was collected while carefully avoiding the buffy-coat. The second centrifugation was conducted for 10 minutes at 4°C and 16,000 g. The supernatant was transferred to 1.5 ml tubes before performing the extraction of cirDNA or being stored at −20°C. CirDNA extraction was performed with the Qiagen Blood Mini kit, following all steps of the protocol. In all, 0.2 to 1 ml of plasma was extracted in several successive passes on a column. The final elution volume was 80 to 130 µl and eluates were frozen at −20°C prior to analysis by qPCR. Freeze-thaw cycles should be avoided to reduce the generation of cirDNA fragmentation and the extracts are not kept longer than 3 months at −20°C.

Q-PCR analysis. CirDNA analysis was performed by a qPCR technique developed in our laboratory, and clinically validated previously. The method is based on an innovative design of short amplicons (60–100 bp) targeting a wild-type sequence of the gene, (here the KRAS nuclear gene and the mitochondrial Cytochrome oxi-
dase III gene, \textit{MT-CO3}). Quantification of this amplicon gives an estimation of the total cirDNA and McirDNA concentration, respectively. For the quantification of cirDNA, we amplified of a 67 bp-length sequence of the KRAS gene with the following primers: forward (5′ CCCTGACATACTCCCAAGGA 3′) and reverse (5′ CCTGGGTTTCAAGTTATATG 3′). For McirDNA, we amplified a 67 bp-length sequence of the cytochrome oxidase sub-unit 3 mitochondrial gene with the following primers: forward (5′ GACCACCAATCACATG 3′) and reverse (5′ TGAAGGCCCCTGTGTA 3′). These primers were designed using the Primer 3 software according to the following requirements: (i) Tc ranging from 50 to 64°C; (ii) GC-content between 40 and 60%; (iii) size from 18 to 23 NT; (iv) amplicon size ranging from 60 to 100 bp. We performed local-alignment analyses with the BLAST program to confirm the specificity of the designed primers. All sequences were checked for self- or inter-molecular annealing with nucleic-acid-folding software (Mfold and oligoAnalyzer 1.2). Oligonucleotides were synthesized and HPLC-purified by Eurofins (Ebersberg, Germany) and quality control of the oligonucleo-
tides was performed by MALDI-TOF. For all analyses, negative controls and standard curves were used. All tests are performed in triplicate with 5 µl of DNA extract in a 25 µl reaction volume, on a CFX96 instrument using CFX manager software (Bio-Rad). This method (qPCR, primer design, program) and technical validation have been described previously. The mCRC patient’s blood samples were excluded if the total cirDNA concentration, due to a problem of pre-analytic treatment or even for unknown reasons, was below a quality threshold.

The DNA concentration quality threshold was 3 ng/mL (about 900 copies/mL) for cancer patients. Note, this value corresponds to about the half of the median concentration found for healthy individuals (N = 104, median NcirDNA concentration = 5.43 ng/mL of plasma corresponding to 1645 copies/mL). In addition to quantifying cirDNA by targeting two different sequences on two different chromosomes, an experiment based on Poisson law distribution showed the accuracy of our cirDNA amount measurement (Supplementary Fig. S8). It should be noted that our Q-PCR systems enable the detection of a single genome copy (Supplementary Table S4), and that we have previously shown that targeting a DNA sequence of the same size or longer than the input DNA fragment produced a similar PCR yield. The measurement of the total cirDNA concentration by targeting a \textit{KRAS} sequence was routinely controlled by quantifying a \textit{BRAF}-based value is 1.3 to 1.8-fold higher than that of the \textit{BRAF}-based value when using the reported Q-PCR primer systems; otherwise, a second analysis is performed. Note, \textit{BRAF} analysis data from 33 healthy individuals were not available. In addition, we excluded 26 patient plasmas (17 mCRC and 9 healthy) in which the total cirDNA concentration, due to a problem of pre-analytic treatment or even for unknown reasons, was below a quality threshold.

CirDNA calibration assay. \textit{NcirDNA}. A genomic DNA extract from human wild-type \textit{KRAS} colorectal cells was used for the NcirDNA calibration assay. Initial genomic DNA solution concentration and purity were determined by measuring optic density at λ = 260 nm, 230 nm and 280 nm, with an Eppendorf BioPhotometer\textsuperscript{®} D30. Starting genomic DNA concentration was adjusted to 1800 pg/µl for the first dilution point, according to optic density measurement at λ = 260 nm. A qPCR standard curve was obtained by 6 successive dilutions of the vector solution (1800, 180, 45, 20, 10 and 5 pg/µl). The standard curve was used to determine the NcirDNA concentra-
tion of the mCRC patients and healthy individuals and calculate the NcirDNA copy number per milliliter of plasma.

\textit{McirDNA}. A 3382-pb human ORF vector with a 786-pb \textit{MT-CO3} insert was obtained from ABM good\textsuperscript{®} (accession no.YP_003024032) and used for the McirDNA calibration assay. Initial vector solution concentration...
and purity were determined by measuring optic density at λ = 260 nm, 230 nm and 280 nm, with an Eppendorf BioPhotometer® D30. Starting vector concentration was adjusted at 1800 pg/µl for the first dilution point, according to optic density measurement at λ = 260 nm. A qPCR standard curve was obtained by 6 successive dilutions of the vector solution (1800, 180, 45, 20, 10 and 5 pg/µl). The standard curve was used to determine the McirDNA concentration of the mCRC patients and healthy individuals and calculate the McirDNA copy number per milliliter of plasma.

**CirDNA copy number calculation.**  
\[ Q_{nuclear} = \left( \frac{c}{3} \right) \times \left( \frac{V_{elution}}{V_{plasma}} \right) \]

\( Q_{nuclear} \) is the NcirDNA copy number per milliliter, \( c \) is the NcirDNA concentration (pg/µl) determined by qPCR targeting the nuclear KRAS gene sequence and 3.3 pg is the human haploid genome mass. \( V_{elution} \) is the volume of cirDNA extract (µl) and \( V_{plasma} \) is the volume of plasma used for the extraction (ml).

**McirDNA.**  
\[ Q_{mito} = \left( \frac{c \times Na}{2 \times MW \times L_{vector}} \right) \times \left( \frac{V_{elution}}{V_{plasma}} \right) \]

\( Q_{mito} \) is the McirDNA copy number per milliliter, \( c \) is the McirDNA mass concentration (g/µl) determined by a qPCR targeting the mitochondrial MT-CO3 gene. \( Na \) is Avogadro’s number (6.02\( \times \) 10\(^{23} \) molecules per mole), \( L_{vector} \) is the plasmid length (nucleotides) and MW is the molecular weight of one nucleotide (g/mol). \( V_{elution} \) is the elution volume of cirDNA extract (µl) and \( V_{plasma} \) is the volume of plasma used for the extraction (ml).

**Statistical analysis.**  
For continuous variables, median and range were computed. To investigate their associations with the biologic parameters, univariate statistical analyses were performed using Mann-Whitney U test or Kruskal-Wallis rank test for continuous variables. Moreover, multivariate analyses were carried out using logistic regressions, with a stepwise selection procedure, to investigate known predictive. Odds ratio (OR) with 95% confidence intervals (CIs) are presented. The power of analysis was reduced due to all patients did not have measurements for all variables. All P values reported are two sided. A probability of ≤0.05 was considered to be statistically significant; *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001. Statistical analysis was performed using the STATA 13.1 software (Stata Corporation, College Station, TX).

**Data Availability**  
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

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**Author Contributions**

R.M. and A.R.T. designed the study, developed the methodology, analyzed the data and prepared the manuscript. R.M., Z.A.A.D., A.O., R.T., B.P., C.S., J.A., G.T., S.A. and S.E.M. realized the experiments. R.M., S.T. and C.M. performed the statistical analysis. All of the authors (R.M., Z.A.A.D., A.O., R.T., B.P., C.S., J.A., G.T., S.A., C.M., A.A., S.E.M., P.B. and A.R.T.) discussed the results and approved the manuscript.

**Additional Information**

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