C and 15N natural isotope abundance reflects breast cancer cell metabolism

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Breast cancer is the most common cancer in women worldwide. Despite the information provided by anatomopathological assessment and molecular markers (such as receptor expression ER, PR, HER2), breast cancer therapies and prognostics depend on the metabolic properties of tumor cells. However, metabolomics have not provided a robust and congruent biomarker yet, likely because individual metabolite contents are insufficient to encapsulate all of the alterations in metabolic fluxes. Here, we took advantage of natural 13C and 15N isotope abundance to show there are isotopic differences between healthy and cancer biopsy tissues or between healthy and malignant cultured cell lines. Isotope mass balance further suggests that these differences are mostly related to lipid metabolism, anaplerosis and urea cycle, three pathways known to be impacted in malignant cells. Our results demonstrate that the isotope signature is a good descriptor of metabolism since it integrates modifications in C partitioning and N excretion altogether. Our present study is thus a starting point to possible clinical applications such as patient screening and biopsy characterization in every cancer that is associated with metabolic changes.

Medical applications of stable isotopes are now widespread, like the well-known 13C-urea breath assay for ulcer detection\textsuperscript{1}. This takes advantage of 13C-labelling and thus usually neglects differences in reaction rates between isotopic forms, because the isotopic signal used for diagnosing is far above small natural variations in 13C. By contrast, the use of isotopes at natural abundance exploits such subtle differences (referred to as isotope effects) to identify bottlenecks in metabolic pathways (rate-limiting steps) or the contribution of multiple elemental sources (mass balance), without the need to introduce expensive isotope tracers into the patient. Isotope effects in metabolism are mostly caused by enzymatic reactions that preferentially consume substrates containing either the light or the heavy isotope (isotopologues) and therefore, the natural isotope abundance in metabolites depends on metabolic fluxes and source substrates\textsuperscript{2}. For example, the natural 13C abundance in respired CO2 has been used to trace diet composition and substrate changes during exercise\textsuperscript{3,4}. In cancer biology, the use of natural variations in Cu and S stable isotopes in hepatocellular carcinoma has been attempted recently\textsuperscript{5}. But to our knowledge, no study has looked at alterations of natural isotope abundance in breast cancer. Due to changes in primary C and N metabolism such as increased glycolysis, glutaminolysis and nucleotide synthesis\textsuperscript{6}, important changes in 13C and 15N natural abundance can be anticipated.

To address this question, we examined the isotopic signature of intact breast cancer biopsies (mostly from invasive ductal carcinoma, IDC) and cultured breast cancer cell lines (Supplementary Tables S1 and S2) using elemental analysis coupled to isotope ratio mass spectrometry (EA-IRMS). This technique has been recently shown to be applicable to the biochemical analysis of cancerous cell lines\textsuperscript{7}.

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Results

Isotope signature of breast cancer biopsies. Natural $^{13}$C and $^{15}$N abundance was found to discriminate normal and cancerous biopsies, the latter being significantly $^{13}$C-enriched by ~4‰ and tended to be (though not significantly) $^{15}$N-depleted (Fig. 1a,b). Triple negative tissues (ER- PR- HER2-) tended to be relatively $^{15}$N-depleted (Fig. 1a and Supplementary Table S1). The observed isotopic difference between cancerous and non-cancerous tissues can have been caused by either variation in chemical composition or isotope fractionations in metabolism.

First, there were alterations in total lipid content, which correlated to the $^{13}$C-abundance (Fig. 1c and Supplementary Figure S1). In fact, lipids are typically $^{13}$C-depleted components of human tissues\(^8\) and a systematic $^{13}$C-depletion has been observed in adipose tissue\(^9\). Here, cancerous tissues are on average less lipid-rich, reflecting the lower abundance of adipose cells in breast IDC\(^10\). Cancerous tissues were also enriched in nitrogenous compounds, as shown by the higher N elemental content (Fig. 1d). Regardless of receptor expression, cancerous tissues had a higher elemental content in N ($P<0.05$). In (a,c), each datum is the average of 3 sub-samples. In (d), lower case letters stand for statistical classes, $P<0.05$.
Metabolic alterations in breast cancer biopsies. Targeted metabolomics analyses by GC-MS were then carried out to identify major metabolic alterations. Normal tissues were clearly differentiated from cancerous tissues in both unsupervised (PCA) and supervised (orthogonal partial least square-discriminant analysis, OPLS-DA) analysis (Supplementary Figures S3 and S4), with ethanolamine, methionine and arginine having the highest weight in such a differentiation (Supplementary Figures S3 and S4). Other discriminating metabolites included carbohydrates (such as fructose) and putrescine, and these metabolites have also been found to be significant elsewhere16,17. While the effect on free ethanolamine reflected the general reorchestration of lipid metabolism18, the simultaneous effect on arginine and putrescine strongly suggests perturbations in arginine metabolism (urea cycle), thereby causing changes in N excretion and thus, by mass balance, in 15N-abundance and N elemental content.

Isotopic signature of cell lines. Potentially, the isotopic signature in biopsies may have been influenced by non-controlled epidemiological and clinical factors (e.g., patient age, pharmacological treatments, etc.). Therefore, we also examined different cell lines cultured under isotopically-controlled conditions: one immortalized, non-cancerous cell line (MCF10A) and six breast cancer cell lines (ZR75-1, MCF7, SKBR3, MDA-MB-231, MDA-MB-468 and Cal51, see Supplementary Table S2). The use of cultured cells further allowed us to carry out isotopic compound-specific analyses as well as targeted gene-expression, metabolomics and EA-IRMS analyses (see Methods) on the same samples.

With the exception of ZR75-1 (non-pleural origin, ER+), all other breast cancer cell lines were 15N-depleted (Fig. 2a). The 15N-abundance in DNA, proteins and total soluble metabolites paralleled that in total organic matter (Supplementary Figure S5). The 13C-abundance in non-cancerous cells was found to be lower than in cancerous cells (Fig. 2a). It did not correlate to the lipid content but rather to the 13C-abundance in lipids and loosely, to the content in C4 organic acids (Fig. 2b) which are known to be 13C-enriched19.

Primary metabolism in cell lines. The endometabolome (i.e., intracellular metabolome) was analyzed to gain information on primary C and N metabolism (summarized in Supplementary Figure S6a). Some metabolites such as arginine, urea and glucose were significantly different between cancer and normal cell lines. Close correlations or anti-correlations were obtained between 13C- or 15N-abundance and key metabolites or metabolic ratios such as urea/arginine and fumarate/aspartate (Fig. 3a). As such, cancer cell lines could be differentiated from the MCF10A non-cancerous cells using these ratios (Fig. 3a).

The present results on both metabolome and isotopes suggest that three metabolic pathways are at the origin of the isotopic differences between cell lines: the urea cycle (arginine metabolism), the anaplerotic carbon fixation, and lipid synthesis. Transcript levels of a selection of genes encoding key enzymes of these pathways was examined by qPCR and in most cases, these genes were less expressed in cancer cell lines (Fig. 2c). Thus, rather than substantial changes in enzyme abundance, the alteration of isotope abundance was caused by changes in metabolic fluxes and/or net exchange of metabolites (absorption, excretion and accumulation) with the extracellular medium.

Isotopic efflux in cell lines. We then examined the metabolic composition of the extracellular medium (exometabolome, Supplementary Figure S6b) to determine the preferential distribution (extracellular vs. intracellular) of metabolites and compute isotopic fluxes (isofluxes). The extracellular-to-intracellular difference and
and 14N/15N isotope effects in both deamidation and the urea cycle yield 15N-depleted arginine. The efflux (excreted N) was 15N-depleted and Arg was even more 15N-depleted in cancerous cell lines by up to $\approx 7\%$ (d). Therefore, the relative isotopic efflux represented by Arg excretion (i.e., the quotient of Arg isoefflux to total isoefflux) was large in ZR75-1 and MCF10A and low in other cell lines (e). Asterisks: significant differences between cell lines (*$P < 0.05$; **$P < 0.01$); dashed line: $\delta^{15}N$ of source Gln (d); lower case letters stand for statistical classes ($P < 0.05$) (e). EA, ethanolamine, 2HG, 2-hydroxyglutarate; PGA, phosphoglycerate; PHB, parahydroxybenzoate.

**Discussion**

We showed isotopic differences between healthy and cancer tissues or cells, and provided evidence that these differences are related to urea, anaplerotic C fixation and/or lipid metabolism, and that the balance in nitrogen metabolism is modified in breast adenocarcinoma tissues or cells compared to normal breast tissues or cells thereby altering the natural $^{15}N$-abundance (Fig. 4). The consumption of glutamine (glutaminolysis) is a major pathway in cancerous cells that involves deamidation by glutaminase, thereby producing ammonia which is partly recycled and excreted in the form of urea. Here, source glutamine is naturally $^{15}N$-depleted ($\delta^{15}N = -6.5\%$) and $^{14}N/^{15}N$ isotope effects in both deamidation and the urea cycle yield $^{15}N$-depleted arginine. The efflux (excretion) of arginine does not counterbalance the N influx and this imbalance is compensated for by intracellular build-up and recycling into polyamines and $\beta$-alanine (Fig. 4) thereby causing a general $^{15}N$-depletion of organic matter. In addition to acid metabolism ($C_{4}$-acids and lactate, Fig. 2b, Supplementary Table S3), the urea cycle incorporates bicarbonate (naturally $^{15}C$-enriched) via the action of carbamoyl phosphate synthase and thus, arginine build-up contributes to the enrichment of $^{13}C$ in cells. This appears indeed to be the best explanation considering known isotope effects in metabolism (Supplementary Table S4). In fact, the natural $^{13}C$-enrichment cannot come from a change in total lipid content (as in tumors) since lipid content variation was small (Supplementary Figure S7) and furthermore, the isotope composition in lipids followed that in total organic matter (Fig. 2b).

Quite noticeably, the isotopic pattern seen in tumor biopsies was similar to that found in cell lines, that is, with a tendency to be $^{15}N$-depleted and $^{13}C$-enriched (Fig. 1a). This pattern is unlikely to come from an effect of nutrition, but rather explained by the specific primary C and N metabolism in tumor tissues, as explained above (Fig. 1). The metabolic origin of the $^{13}C$-enrichment was different to that in cultured cells (a decrease in total lipid content in tumors, Supplementary Figure S7 and refs 18,30). This is not surprising since cultured cell metabolism is different from that of cells in situ. By contrast, the mechanism explaining the lower $^{15}N$-abundance was similar: a net influx of $^{15}N$-depleted nitrogen associated with any metabolism. In cancerous cultured cells, there is an increased activity of the urea cycle and non-quantitative excretion of $^{15}N$-depleted arginine. In tumors in situ, it is believed...
that cells are arginine-dependent to sustain polyamine metabolism\textsuperscript{31}, and the auxotrophy of tumors for arginine has been exploited in the development of anticancer therapies\textsuperscript{32}. Therefore cells capture naturally \textsuperscript{15}N-depleted arginine from their microenvironment, thereby leading to a \textsuperscript{15}N-depletion of cellular organic matter. Such a mechanism agrees with the well-known increase in polyamine metabolism\textsuperscript{33} and the fact that urea and glutamine (glutaminase activity) are amongst the best urinary and tumor biomarkers for breast cancer, respectively\textsuperscript{34,35}. Also, differential expression of arginase, a key enzyme of the urea cycle, has been reported in breast cancer cell lines\textsuperscript{36,37}.

The natural abundance in \textsuperscript{13}C and \textsuperscript{15}N thus reflects primary metabolism of cancer cells and thus, offers potential opportunities for development of biomarkers for clinical research. Consistently, healthy and cancerous samples were found to fall in corresponding regions of the isotopic landscape (Fig. 1a), and normal and cancerous cell lines could be differentiated (Fig. 2a). The ability of natural \textsuperscript{13}C- or \textsuperscript{15}N-abundance to further differentiate tumor grade and severity remains to be explored. In addition, there is a potential use the natural isotopic abundance to distinguish between different breast cancer subtypes, as they have been shown to have different metabolic phenotypes\textsuperscript{38}. Also, differential expression of arginase, a key enzyme of the urea cycle, has been reported in breast cancer cell lines\textsuperscript{36,37}.

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Methods

Human Tissues. Fresh human mammary samples were obtained from chemotherapy naïve patients with invasive carcinoma after surgical resection at the Institut de Cancérologie de l’Ouest – Rene Gauducheau, Nantes, France. Informed consent was obtained from patients to use their surgical specimens and clinicopathological data for research purposes, as required by the French Committee for the Protection of Human Subjects (CCPPRB). Ouest IV – Nantes CCPPRB approved use of tumor tissues for this study (6 May 2013: no. 357/2013). The collection of tumors was approved by the French Minister of higher education and research (no. AC-2008-141). This study did not require additional ethical approval. Amongst 17 breast cancer patients, 12 breast cancer samples and 5 pairs of breast cancer and adjacent non-cancerous tissues were obtained. One normal breast sample was obtained from reduction mammoplasty. For tumors, hormonal receptors, histological type and grade information were available, except that a paired of tissue #22/#19 (Supplementary Table S1, u and u’ in Fig. 1a) did not have receptor status available.

Cultured cells. Six human breast cancer cell lines and one non-cancerous cell line (Supplementary Table S2) were used and grown as monolayer cultures (\texttimes10\textsuperscript{7} cells/75 cm\textsuperscript{2} flask). The cancer cells were grown in no-glucose...
no-glutamine DMEM supplemented with 10% FBS and 1% 10,000 units/ml penicillin, 10,000 units/ml streptomycin at 37°C in a humidified atmosphere of 5% CO₂. The non-cancerous cell line (MCF10A) was grown in DMEM/F-12 (no-glucose no-glutamine) supplemented with 5% horse serum, 0.01% EGF (Epidermal Growth Factor), 0.005% cholera toxin, 0.1% insulin, 0.005% hydrocortisone, and 1% HEPES. L-glutamine and D-glucose (4 and 25 mM, respectively) were added separately to culture mediums to ensure these nutrients were isotopeically identical for all cells. The same cells were grown several times to test the biological variability. As soon as cells were confluent (~10⁶ cells), cell counting and extraction were done using triplicate cell culture flasks. One flask was used for cell counting, after harvesting with trypsin, using trypsin blue staining and microscopic hemocytometry. For other analyses carried out on the other flasks, cells were then prepared as an aqueous (isotopes) or methanolic (metabolomics) extract without using trypsin as explained below.

**Extraction and separation of studied fractions.** Human tissues. Human fresh tissues were lyophilised. Lipids were extracted from the dried human tissues (10 mg) with hexane (1 mL). After centrifugation (4000 rpm, 15 min, 4°C), the pellet was kept for metabolomics and the lipids were recovered by evaporation of hexane under a stream of N₂ gas at 40°C.

**Cultured cells.** After culture, cells were washed twice with PBS, and 3 mL of cold water was added to the flask. Cells were detached from the flask using a cell scraper. The lysed cell preparation was kept at −20°C for 48 h and then lyophilized. Different fractions (lipids, total soluble fraction (TSF), protein and DNA) were extracted and separated from the dry cell material. To obtain a lipid fraction, lyophilized cells (40 mg) were ground and extracted twice under agitation with 2 mL cyclohexane for 5 min at 20°C. After centrifugation (4000 rpm, 5 min, 20°C), the lipid fraction (4 mL of cyclohexane) was dried under a stream of N₂ gas at 40°C and the dry residue stored at −20°C until required. To obtain a total soluble fraction, the pellet (insoluble in cyclohexane) obtained after centrifugation were ground and extracted twice with 2 mL methanol/water (80:20 v/v) for 5 min at 20°C. After centrifugation (4000 rpm, 5 min, 20°C), the total soluble fraction was dried under a stream of N₂ gas at 40°C and the dry residue stored at −20°C until required. The pellet (insoluble in methanol/water) consisted of a protein fraction.

For DNA purification, lyophilized cells (15 mg) were reconstituted with 1 mL water and extracted with an equal volume of phenol/chloroform in an ultrasonic bath for 30 min. After centrifugation (4000 rpm, 5 min, 4°C), the aqueous phase was filtered to remove proteins and the DNA was precipitated with an equal volume of phenol/chloroform in an ultrasound bath for 30 min. After centrifugation (4000 rpm, 5 min, 20°C), the total soluble fraction was dried under a stream of N₂ gas at 40°C and the DNA purified by ethanol (70%) to remove impurities, and dried under a stream of N₂ gas at 40°C.

**13C and 15N abundance by elemental-analysis coupled to isotope ratio mass spectrometry (EA-IRMS).** Subcellular fractions (see above) were lyophilized for EA-IRMS analysis, in order to detect their 13C and 15N isotopes. From each sample ~0.7 mg was weighed with 10⁻² g precision balance (Ohaus Discovery DV215CD, Pine Brook, New Jersey, USA) into each of three tin capsules (solds “light” 5 × 9 mm, Thermo Fisher Scientific, www.thermo.com) for isotope analysis. EA-IRMS analyses were performed as previously described. The 13C and 15N abundance was first expressed as δ values (δ¹³C and δ¹⁵N, in ‰) as δ = R/Rs − 1 where R is the heavy-to-light isotope ratio and Rₛ stands for the isotope ratio in the international reference (V-PDB for 1³C and air N₄ for ¹⁵N).

The isotope abundance in sample fractions was expressed relative to that in total organic matter using the enrichment (in ‰) e = δœur − δₑₑₑ wherein δₑₑₑ is the δ value in the fraction of interest (lipids, proteins, etc.) and δₑₑₑ the (non-corrected) δ value in total organic matter. The enrichment e is then positive if the sample fraction is enriched in the heavy isotope and negative if it is depleted. In this way, e nearly equals the opposite of the isotope fractionation Δ (which is given by [δₑₑₑ − δₑₑₑ]/[δₑₑₑ + 1] ≈ δₑₑₑ − δₑₑₑ). For total organic matter in Figs 1 and 2 and Supplementary Figure S3, δ values were corrected to allow comparisons between patients (that may have different nutritional preferences) and between cell lines (that had slightly different culture media), as follows:

In cultured cells, it was corrected for the δ value of the elemental C and N source using the classical relationships accounting for isotope fractionation: δₑₑₑ = (δₑₑₑ − Δ)/(Δ + 1) where Δ = δₑₑₑ − δₑₑₑ and δₑₑₑ is the δ value of the elemental source and δₑₑₑ is δ value used as the reference (here, the average of δₑₑₑ values was used as the reference value; note that any change in this value would not change the pattern seen here, it would only change the absolute scale of δₑₑₑ values). δₑₑₑ was the δ¹⁵N of and δ¹³C of source amino acids and sugars, respectively, in intact culture medium. When expressed as a fractionation (Fig. 2a), the isotope composition in total organic matter of cells is given as e = δ − δₑₑₑ. That way, e values are positive when it is enriched in the heavy isotope and negative when it is depleted, as for e in individual cell fractions (see above).

For biopsies, δₑₑₑ could not, quite obviously, be known because food composition of patients was not controlled. We therefore had to use a correction that could be calculated independently of food composition. The major influence of nutrition on elemental composition and isotopes is the protein/fat ratio of ingested food, with fat-rich and protein-poor food causing a general increase in %C (and decrease in %N) and a general decrease in both δ¹³C and δ¹⁵N in body tissues. Therefore, we renormalized the δ values in total organic matter by taking into account the %C and %N using δₑₑₑ = δ(1 − f)/τ where f is the elemental composition relative to the average and τ is the apparent isotope fractionation between the whole population δ value (average isotope composition of human tissues in European western (French) population (b), −22‰ and 9‰ for ¹³C and ¹⁵N, respectively) and the average of our samples (bₑₑₑ): τ = bₑₑₑ − bₑₑₑ. For example, in this way, protein-rich samples (lean patients) that are naturally impoverished in elemental C and richer in ¹⁵N will have a negative correction for δ¹³C. Of course, such a correction was not ideal because the knowledge of the average isotope composition of ingested food would have been better. In practice, this correction was minimal (on average, 0.02‰ for ¹³C and 0.5‰ for ¹⁵N) and did not cause a substantial bias in our data.
\(^{13}\)N-abundance and amino acid content using gas chromatography - combustion - isotope ratio mass spectrometry (GC-C-IRMS). When cells were confluent, the culture medium was kept (10 mL from \(~10^7\) cells), and cells (~10^8) were washed with PBS and then lysed in 36 mL of 80% methanol. After homogenization, methanolic extracts were dried under a stream of \(N_2\) gas at 40°C. Amino acids were prepared as their N-pivaloyl, O-isopropyl esters for GC-C-IRMS analysis, and used for both quantification (\(\mu\)mol) and \(^{13}\)N values (%) as previously described^40.

\(^{13}\)C-abundance in metabolites using liquid chromatography coupled to chemical oxidation and isotope ratio mass spectrometry (LC-co-IRMS). For LC-co-IRMS analysis, \(~10^7\) cells were used in 12 mL of 80% methanol. LC-co-IRMS analysis was performed as previously described, using water as a mobile phase and a RS Pak KC-811 column (Shodex)^41. \(^{613}\)C values were corrected using pure standard solutions at different concentrations. In practice, such corrections due to non-linearity were very small, about 0.2‰.

**GC-MS metabolomics. Human biopsies.** If material remained after lyophilization and lipid extraction, an additional \(~10–15\) mg of tissue was ground and dissolved in a standard solution of DL-2-Aminobutyric acid (0.2 mM, 950 \(\mu\)L) and methanol/water (950 \(\mu\)L, 80:20 v/v) was added to extract the soluble metabolites. After centrifugation (14000 rpm, 15 min, 4°C), the soluble fraction was mixed with standard solution of ribitol (2 mM, 95 \(\mu\)L). The solution was centrifuged again and three aliquots of 200 \(\mu\)L were dried using a speed-vac before derivatization and GC-MS analysis as described previously^41,42.

**Cultured cells and extracellular medium.** About \(~5 \times 10^7\) cells were dissolved in a solution of methanol/water (7 mL, 60:40 v/v) containing 500 \(\mu\)L of a standard solution DL-2-aminobutyric acid (0.2 mM). After centrifugation (14000 rpm, 15 min, 4°C), the soluble fraction was mixed with another standard solution of ribitol (2 mM, 50 \(\mu\)L). The solution was centrifuged again and three aliquots of 1.4 mL, i.e. 20% of \(~5 \times 10^7\) cells, were dried using a speed-vac before derivatization and GC-MS analysis. For extracellular medium, the same standard solutions [D,L-2-aminobutyric acid (0.2 mM, 500 \(\mu\)L) and ribitol (2 mM, 50 \(\mu\)L) and methanol/water solution (500 \(\mu\)L, 60:40 v/v)] were added to 500 \(\mu\)L of media collected from the cultured cells. After a centrifugation, three aliquots of 300 \(\mu\)L were prepared for GC-MS analysis. All metabolomics data were normalized to the specific number of cells so that the exo- and endo-metabolome could be directly compared.

**RT-qPCR.** RNA was isolated from cell lines and 500 ng RNA was reverse transcribed as previously described^43. Quantitative PCR was done using the Maxima SYBR Green/ROX qPCR Master Mix (Life Technologies) and the MX4000 instrument (Stratagene, Basel, Switzerland), according to the manufacturer’s instructions. The endogenous housekeeping genes RPLPO, HPRT1 and ACTB1 were used for normalization. Relative quantification was carried out by using the \(\Delta\Delta\)CT method. Primer sequences are listed in Supplementary Table S5.

**Statistics.** Pair-wise comparisons were carried out using Student-Welsh tests, with a p-threshold value of 0.01 unless otherwise stated in text. Metabolomics were examined using unsupervised analysis with principal component analysis (PCA) or supervised analysis with orthogonal partial least square discriminant analysis (OPLS-DA), both carried out with Simca (Umetrics). Heat maps and hierarchical clustering (HCL) were done using the open source software MeV^44. The HCL was carried out using the cosine correlation as a measure of proximity.

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Author Contributions

I.T. and G.T. designed and supervised the project. E.M., I.A. and C.M. collected and generated the isotopic data and performed isotopic analyses; J.L. performed qPCR analyses; Metabolomics data were performed by F.G. and S.B.-N. provided tissue and cell samples. S.B.-N. and A.C.B. advised on breast cancer metabolism context. G.T. and I.T. performed statistical analyses, prepared the figures and tables and wrote the manuscript; S.B.-N. and A.C.B. revised the manuscript; all authors viewed, had the opportunity to comment on and approved the final draft.

Additional Information

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