Bonded Cumomer Analysis of Human Melanoma Metabolism Monitored by 13C NMR Spectroscopy of Perfused Tumor Cells.

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Bonded Cumomer Analysis of Human Melanoma Metabolism Monitored by $^{13}$C NMR Spectroscopy of Perfused Tumor Cells*

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A network model for the determination of tumor metabolic fluxes from $^{13}$C NMR kinetic isotopomer data has been developed and validated with perfused human DB-1 melanoma cells carrying the BRAF V600E mutation, which promotes oxidative metabolism. The model generated in the bonded cumomer formalism describes key pathways of tumor intermediary metabolism and yields dynamic curves for positional isotopic enrichment and spin-spin multiplets. Cells attached to microcarrier beads were perfused with 26 mM [1,6-$^{13}$C$_2$]glucose under normoxic conditions at 37 °C and monitored by $^{13}$C NMR spectroscopy. Excellent agreement between model-predicted and experimentally measured values of the rates of oxygen and glucose consumption, lactate production, and glutamate pool size validated the model. ATP production by glycolytic and oxidative metabolism were compared under hyperglycemic normoxic conditions; 51% of the energy came from oxidative phosphorylation and 49% came from glycolysis. Even though the rate of glutamine uptake was ~50% of the tricarboxylic acid cycle flux, the rate of ATP production from glutamine was essentially zero (no glutaminolysis). De novo fatty acid production was ~6% of the tricarboxylic acid cycle flux. The oxidative pentose phosphate pathway flux was 3.6% of glycolysis, and three non-oxidative pentose phosphate pathway exchange fluxes were calculated. Mass spectrometry was then used to compare fluxes through various pathways under hyperglycemic (26 mM) and euglycemic (5 mM) conditions. Under euglycemic conditions glutamine uptake doubled, but ATP production from glutamine did not significantly change. A new parameter measuring the Warburg effect (the ratio of lactate production flux to pyruvate influx through the mitochondrial pyruvate carrier) was calculated to be 21, close to upper limit of oxidative metabolism.

Cancer cells exhibit metabolic patterns that differ significantly from those of terminally differentiated adult tissues. They require rapid production of both energy and biosynthetic precursors to sustain a high rate of proliferation. The network of biochemical pathways underlying these processes is complex. Most of the individual enzymatic conversions involved in this network have probably already been identified, but a comprehensive and quantitative description of the fluxes involved is lacking. Methods to produce such descriptions will likely be very useful in the design of new therapeutics that specifically target the metabolic abnormalities of cancer as well as in clinical implementation of non-invasive methods for detection of cancer and its therapeutic response.

The kinetics of $^{13}$C isotope labeling as monitored by NMR spectroscopy has been used to measure flux through various metabolic pathways of perfused isolated cells and organs and in vivo (1–4). Kinetic data provide estimates of absolute flux, whereas steady-state data measure relative flux. Both types of analysis require validated mathematical models, which have been formulated for the heart (5–10), liver (11–13), brain (14–16), and tumors (17). Only the heart model has been validated by comparison between model-derived and experimental parameters (6, 7). Modeling of tumor metabolic fluxes has been limited to steady-state analysis with dynamic measurements of glucose uptake and lactate production subsequently added to convert to absolute fluxes; none of these models has been validated. To the best of our knowledge, this is the first validated dynamic flux model of tumor intermediary metabolism based on $^{13}$C NMR isotope kinetic data.

Mathematical models of $^{13}$C magnetic resonance spectroscopy data are based on measurement of positional isotopic enrichment and/or on isotopomer analysis, which takes into account spin-spin coupling of covalently bonded labeled carbon atoms. Bonded cumomer analysis includes both positional enrichment and isotopomer modeling and facilitates the formulation of kinetic differential equations. This method requires fewer equations than classical isotopomer analysis and is based on internal symmetry properties of metabolite isotopomers.
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Our laboratories have developed a Continuous Stirred Tank Bioreactor suitable for high resolution 13C NMR spectroscopy of cultured cancer cells that can produce detailed spectra with good temporal resolution while simultaneously monitoring the oxygen consumption rate (18–21). Specifically, we measured the time course for labeling of key intermediates such as glutamate, aspartate, alanine, lactate, and fatty acyl groups in real time during growth with 13C-labeled glucose and unlabeled glutamine as substrates.

Here, we demonstrate that absolute flux measurements of perfused tumor cells can be obtained in several hours from ∼5 × 10⁶ tumor cells in this bioreactor system, and the data can be analyzed using a refined isotopomer analysis method called bonded cumomer (BC) analysis (22) that utilizes only ∼210 differential equations to describe most of tumor intermediary metabolism, whereas the heart model of Chance et al. (5) required ∼325 differential equations to describe just TCA cycle activity in terms of positional enrichment. To validate the BC model, we compared model-derived and experimental values of the following parameters: O2 consumption rate (MRO2), glutamate pool sizes, as well as rates of glucose consumption and lactate release. This study permits direct measurements of energy production by glycolytic and mitochondrial pathways and quantitative evaluation of the Warburg effect. It also provides a basis for future studies of the effects of various therapeutic interventions on tumor metabolism. Whereas studies of perfused tumor cells in a bioreactor facilitate accurate and reproducible measurements, this model is also applicable to in vivo studies in animal models and humans. Recent studies of ex vivo surgically resected human brain (23) and lung (24) tumors point to the potential utility of in vivo 13C MRS clinical applications.

Experimental Procedures

Cell Culture—DB-1 human melanoma cells were derived from a lymph node metastasis (Thomas Jefferson University Hospital, Philadelphia, PA). The cells were obtained before administration of any treatment. Cells were cultured from the tumor and cryopreserved after the 16th passage. The presence of melanoma cell surface antigens was confirmed with monoclonal antibodies (25). For routine culture, DB-1 cells were grown as monolayers at 37 °C in 5% CO2 in α-minimal essential medium supplemented with 10% fetal bovine serum, 2 mM glutamine, 26 mM glucose, and 1% (v/v) non-essential amino acids and 10 mM HEPES buffer. The doubling time in tissue culture flasks was 48 h (26).

Cell Perfusion and NMR Spectroscopy—For cell perfusion studies, DB-1 cells were cultured in DMEM with 1% (v/v) non-essential amino acids (Invitrogen), 10% fetal bovine serum, 10 mm HEPES, 4 mM glutamine, and 26 mM glucose (complete DMEM). During NMR experiments, ∼5 × 10⁶ cells were grown on the surface of non-porous microcarriers that had a mean diameter of 170 μm (SoluHill, Ann Arbor, MI). The surface of the microcarriers was coated with either collagen or ProNectinF (20) to enhance cell attachment and proliferation. The microcarriers were tightly packed inside a 20-mm NMR tube. They were perfused continuously with the system shown in Fig. 1. A detailed description of the perfusion system, including flow rates, cell adhesion procedures, etc., has been presented previously (20).

A peristaltic pump (Masterflex, Cole Parmer, Chicago) was used to circulate medium through the microcarriers at a flow rate of 12 ml/min. Before entering the cell mass, the medium flowed through a gas exchange module where carbon dioxide was removed, and oxygen was added across a silicone membrane (thin-wall silicone tubing). Subsequently, the medium was warmed to 40 °C, and the oxygen level was measured continuously with a polarographic oxygen probe (Mettler-Toledo, Columbus, OH). The steady-state oxygen level of the medium at the inlet of the cell mass was maintained near air saturation. A second polarographic oxygen probe was used to detect the oxygen level of the medium coming out of the NMR tube so that the oxygen consumption rate of the culture could be determined continuously. A pH probe (Mettler-Toledo, Columbus, OH) was also used downstream of the outlet oxygen probe. The pH of the medium was controlled by adjusting the level of CO2 in the gas exchange module. The temperature of the medium entering the NMR tube was fine-tuned with a microstat-controller and an electrical resistance heater to 37 ± 0.2 °C. The glucose concentration in the recirculating medium was maintained at a constant level (clamped) by continuously feeding fresh medium and removing depleted medium from a recirculation bottle. The flow rate of fresh medium to the system was typically 24 ml/h, and the volume of recirculating medium was 120 ml. During the 13C experiments, 32 mM [1,6-13C2]glucose was introduced into the feed to the system, and the recirculating glucose level wasclamped at 26 mM by adjusting the feed rate.

13C NMR spectra were acquired with standard 1H decoupled NOE 13C acquisition on a 9.4 tesla/89 mm vertical bore Varian spectrometer (Varian Inc., Palo Alto, CA). The acquisition parameters were as follows: 60° pulse angle, 1.2-s repetition rate, 25,000 Hz spectral width, 16,384 points, and 750 transients per spectrum. To ensure that 1H decoupling did not raise the temperature of the perfusate above 37.5 °C, the temperature of the medium leaving the NMR tube was continuously monitored with a thermocouple. Free induction decays were apodized by exponential multiplication (2 Hz) for signal to noise enhancement. Nuts NMR (Acorn NMR, Fremont, CA) was used to determine resonance peak areas. The cell number in the NMR tube was determined from the total NTP level as described previously (27). 31P spectral parameters were 60° pulse angle, 1-s repetition rate, 15,000 Hz, and 1200 transients.

In this work, melanoma cells were initially studied under normoxic hyperglycemic conditions to enhance lactic acidosis (28, 29) and subsequently under euglycemic conditions to more closely simulate in vivo conditions. The effects of glucose con-
centrations on flux through various pathways of tumor metabolism were evaluated. In the initial studies, the concentration of glucose in the perfusate was maintained at 26 mM, a value that we believe would be the upper limit for a therapeutic procedure.

In subsequent studies, the glucose concentration was 5 mM. The kinetics of labeling in lactate, glutamate, and other metabolites was monitored over an ~6-h time course. The results were used to estimate intracellular fluxes. The estimated TCA cycle flux was compared with the oxygen consumption flux to test the accuracy of the calculations.

**Pentose Phosphate Pathway (PPP) Labeling**—DB-1 cells were grown in T-150 flasks in α-minimal essential medium. At about 70% confluency of cells in the flask, the medium was replaced with DMEM with 5 mM [1,2-13C]glucose (Cambridge Isotope Laboratories, Inc., Cambridge, MA) and incubated for 8 h at 37 °C and 5% CO2. The cells were harvested by using 0.05% trypsin/EDTA (Invitrogen), rinsed twice with phosphate-buffered saline (PBS) at 4 °C, and frozen. The cells were homogenized after adding 4 ml of 12 M perchloric acid and were spun at 13,000 × g for 30 min. The supernatant was neutralized with 3 M KOH, lyophilized, and dissolved in 0.6 ml of D2O. High resolution NMR spectra of the metabolites were generated using a 9.4-tesla spectrometer (Agilent, Palo Alto, CA) equipped with a 5-mm broadband probe. Proton-decoupled 13C spectra were acquired with the following parameters: sweep width = 20 kHz, number of points = 128,000, flip angle 30°, repetition time = 7.3 s, and number of average = 7000. The proton-observed carbon-edited NMR technique (30) was employed to differentiate between the 13C-labeled and -unlabeled lactate signals in the proton spectra with the following parameters: sweep width = 5 kHz, number of points = 32,000, repetition time = 12 s, echo time = 1/7.8 ms, and number of average = 64. Experiments were run in duplicate. iNMR software (Mestrelab Research, Spain) was used for spectral fitting and integration.

**Glutaminolysis and Fatty Acids Labeling**—DB-1 cells were grown on C104-1521 microcarriers (SoloHill) in uncoated T-75 flasks with α-minimal essential medium. At about 70% confluency of cells on the microcarrier beads, the medium was replaced with DMEM with 5 mM 1,2-13C]-glucose (Cambridge Isotope Laboratories, Inc.), Cambridge, MA) and incubated for 8 h at 37 °C and 5% CO2. The cells were harvested by using 0.05% trypsin/EDTA (Invitrogen), rinsed twice with phosphate-buffered saline (PBS) at 4 °C, and frozen. The cells were homogenized after adding 4 ml of 12 M perchloric acid and were spun at 13,000 × g for 30 min. The supernatant was neutralized with 3 M KOH, lyophilized, and dissolved in 0.6 ml of D2O. High resolution NMR spectra of the metabolites were generated using a 9.4-tesla spectrometer (Agilent, Palo Alto, CA) equipped with a 5-mm broadband probe. Proton-decoupled 13C spectra were acquired with the following parameters: sweep width = 20 kHz, number of points = 128,000, flip angle 30°, repetition time = 7.3 s, and number of average = 7000. The proton-observed carbon-edited NMR technique (30) was employed to differentiate between the 13C-labeled and -unlabeled lactate signals in the proton spectra with the following parameters: sweep width = 5 kHz, number of points = 32,000, repetition time = 12 s, echo time = 1/7.8 ms, and number of average = 64. Experiments were run in duplicate. iNMR software (Mestrelab Research, Spain) was used for spectral fitting and integration.

** bumetabolism**—Glutaminolysis and Fatty Acids Labeling—DB-1 cells were grown on C104-1521 microcarriers (SoloHill) in uncoated T-75 flasks with α-minimal essential medium. At about 70% confluency of cells on the microcarrier beads, the medium was replaced with DMEM with 2 mM l-glutamine (U-13C, 99%) (Cambridge Isotope Laboratories, Inc.). Two concentrations of glucose were used in the medium, 5 and 25 mM. After 8 h of incubation, the cells were carefully stripped from the microcarriers using mechanical agitation. Microcarriers were allowed to settle, and suspension containing cells was collected. Cells were rinsed twice with PBS at 4 °C and frozen until metabolite extraction. For fatty acid labeling, cells were grown, incubated, and harvested as above with the exception that the incubation was performed for 24 h and the label used was l-glutamine (13C, 99%; 15N, 99%) (Cambridge Isotope Laboratories, Inc.).

**LC/MS**—The isotope-enriched cells were washed twice with PBS and resuspended in 700 μl of ice-cold methanol. Samples were pulse-sonicated for 30 s and centrifuged at 16,000 × g for 10 min. The supernatant was then transferred to another tube and evaporated to dryness under nitrogen. Samples were resuspended in 100 μl of water and analyzed using an Agilent 1200 series HPLC system coupled to an Agilent 6460 triple quadrupole mass spectrometer equipped with an ESI source operated in the negative ion mode. The injection volume was 10 μl. Analytes were separated by reverse-phase ion-pairing chromatography utilizing a Phenomenex C18 column (150 × 2.0 mm, 3 μm particle size) at a flow rate of 200 μl/min and maintained at 45 °C. A two-gradient solvent system was used, with solvent A as 400 mM 1,1,1,3,3,3-hexafluoro-2-propanol and 10 mM disopropylethylamine in water, and solvent B as 400 mM 1,1,1,3,3,3-hexafluoro-2-propanol and 10 mM disopropylethylamine in methanol. The linear gradient conditions were as follows: 2% B at 0 min; 2% B at 4 min; 95% B at 9 min; 95% B at 11 min; and 2% B at 11.5 min, followed by a 3.5-min equilibration. The Agilent 6460 mass spectrometer operating conditions were as follows: gas temperature was set at 320 °C, and the gas flow was set to 8 liters/min. The sheath gas temperature was 400 °C, and the sheath gas flow was set to 10 liters/min. The capillary voltage was set to 3000 V, and the nozzle voltage was set to 1000 V.

**Bonded Cumomer Metabolic Network Model**—To calculate fluxes through key pathways of tumor energy metabolism from 13C isotopic labeling data, we have utilized a novel three-compartment metabolic network model with the bionetwork schematically depicted in Fig. 2 and applicable to Continuous Stirred Tank Bioreactor systems. The model includes medium, cytosolic and mitochondrial compartments interconnected with various transporters. The cytosolic membrane contains glucose (GLUT), monocarboxylic acid (MCT), and glutamine (GLN) transporters to the cytosolic compartment. Tumors generally express GLUT1 and GLUT3, but only GLUT1 is currently included in the model; similarly DB-1 contains MCT1 and MCT4 in approximately equal proportions (26), but because the Km value of MCT1 is ~1/10th that of MCT4, we have only included MCT1 in the current model. We have included the GLN transporter (alanine-serine-cysteine (GLN) transporter 2) because of the critical role of glutamine in tumor energetics and metabolism. The BC model can readily be extended to incorporate other isoforms of MCT and GLUT as well as transporters for other amino acids and metabolites.

The mitochondrial pyruvate carrier, which was recently identified in all mammalian cells (31–33), was included to permit access of pyruvate to the PDH complex for conversion to acetyl-CoA and incorporation into the TCA cycle. Pyruvate is also linked to the TCA cycle via pyruvate carboxylase. Citrate, glutamate, and malate are included in the cytosolic compartment with direct links through the mitochondria to the TCA cycle. As already noted, the mitochondrial compartment contains the complete TCA cycle with links to the cytosol for citrate and malate. α-Ketoglutarate is linked to glutamate via glutamate dehydrogenase (GLUD) and two aminotransferases, aspartate aminotransferase (GOT) and alanine aminotransferase (GPT).

A constructed BC dynamic model adapted to perfusion bioreactor experiments was used to fit the 13C time courses of labeled lactate, glutamate, and fatty acids to determine the corresponding metabolic fluxes and the transport parameter for bidirectional lactate transport through the cell membrane. In the model, the perfused [1,6-13C2]glucose and glutamine (labeled in some experiments, see below) are transported from the extracellular medium to the DB-1 cells assuming that reversible non-steady-state Michaelis-Menten facilitative
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transport kinetics occurs through glucose (GLUT family) or glutamine (alanine-serine-cysteine (GLN) transporter 2) transporters as shown in Equation 1,

\[ J_{tr}^{i} = \frac{f_{\text{max}}^i \left( \frac{M_{ei}}{K_{mi}^m} - \frac{M_i}{K_{mi}^m} \right)}{1 + \frac{M_{ei}}{K_{mi}^m} + \frac{M_i}{K_{mi}^m}} \]  

(Eq. 1)

where \( J_{tr}^{i} \) is the net transport flux for boundary species \( i \) between media and cytosol or cytosol and mitochondria; \( M_{ei} \) is the extracellular concentration for species \( i \); \( f_{\text{max}}^{i} \) is the maximal transport rate; \( K_{mi}^m \) is the Michaelis-Menten constant for transport. The labeled lactate from the cell is transported into the medium and obeys the same Michaelis-Menten kinetics through the mono-carboxylic acid transporter (MCT family). In this study, facilitated media-cell transport is assumed for glucose, lactate, and glutamine.

The metabolic network includes glycolysis, TCA cycle, \( \alpha \)-ketoglutarate-glutamate, and oxaloacetate-aspartate exchange through the following: malate-aspartate shuttle; anaplerosis through \( \alpha \) pyruvate carboxylase activity, \( \beta \) at the succinyl-CoA level, and \( \gamma \) glutaminolysis through mitochondrial glutaminases (GLS and GLS2); pyruvate recycling through cytosolic (NADP+-dependent ME1) and mitochondrial malic enzymes (NAD+-dependent ME2 and NADP+-dependent ME3); lactate dehydrogenase activity; and transport processes. In this study, PPP fluxes through oxidative and non-oxidative branches were included in the modeling analysis. The network also includes \textit{de novo} lipogenesis and fatty acid oxidation.

The model was expressed mathematically using two types of mass balance equations, i.e. for chemical and isotope variables. The mass balance equation for the total metabolite concentration in the medium, cytosol and mitochondria is Equation 2,

\[ \frac{dM_i}{dt} = J_{tr}^i + \sum_j v_{ij} F_{ij} - \sum_k v_{ik} F_{ik} \]  

(Eq. 2)

where \( M_i \) is the intracellular concentration of the \( i \) species; \( J_{tr}^i \) is net transport flux for boundary species \( i \) between media and cytosol or cytosol and mitochondria; \( F_{ij} \), and \( F_{ik} \) are normalized reaction fluxes that produce \( j \) or utilize \( k \) cellular species \( i \); and \( v_{ij} \) and \( v_{ik} \) are corresponding stoichiometric coefficients. For extracellular boundary metabolites in the medium, dynamic mass balance has the following general form as shown in Equation 3,

\[ \frac{dM_e}{dt} = r_e \cdot J_{tr}^e \]  

(Eq. 3)

where the extracellular species concentration is \( M_e \) and \( r_e \) is the ratio of cell volume to media volume. An additional term for the medium flow rate was also included. Note that we do not assume here that media boundary metabolites are at constant concentrations, and this assumption (or lack of assumption) is common for bioreactor systems.

Together with the initial metabolome state vector \( M_0 \in R^N \) and fluxome \( F_0 \in R^M \) (subscript 0 refers to baseline steady-state values and reaction rates) isotopomer dynamics of this system are formulated as an initial value problem for ordinary differential equations in terms of bonded cumomer fractions as the state variables. For parallel monomolecular reactions, metabolite \( ^{13}C \) cumomer mass balance was expressed in the form shown in Equation 4,

\[ \frac{d\pi_{ij}}{dt} = \sum_{j} F_{ij} \pi_{ij} - \sum_{k} F_{ik} \pi_{ik} \]  

(Eq. 4)

where metabolite \( M \) is downstream of another metabolite \( S \). The total outflux \( \sum F_{ik} \) balances total influx \( \sum F_{ij} \). \( \pi_{ij} \) represents the total pool size of metabolite \( M \), whereas \( \pi_{ij} \) represent the \( i \)th cumomer fraction of metabolite \( M \) and metabolite \( S \) (i bonded cumomer), respectively.

Isotopomer balance equations were derived for every metabolite bonded cumomer of orders 1, 2, and 3 (e.g. glutamate, glutamine, and aspartate) as described by Shestov et al. (22). This resulted in a set of ~210 differential equations. Fine structure multiplets were completely described by each metabolite's bonded cumomers of order 1, 2, and 3. The use of bonded cumomers leads to a reduced number of equations compared with a model, including all possible isotopomers, while retaining all the NMR-measurable isotopomer information. The term cumomer was first proposed by Wiechert et al. (34), and the concept of cumomer was used in early studies by Muzykantov and Shestov (35). There are connection matrices between the "bonded cumomer" \( \pi \) vectors, which reflect subsets of particularly labeled isotopomers and the vectors of singlets, doublets, triplets, and quartets of metabolite "fine structure" of \( ^{13}C \) NMR spectra. One can derive kinetic equations in the form of fine-structure spectroscopically defined NMR data using matrix connection equations. For example, for a metabolite with five carbon atoms such as glutamate, the equation for carbon 2 would be as shown in Equation 5,

\[ \begin{pmatrix} c_{2q} \\ c_{2d12} \\ c_{2d23} \\ c_{2s} \end{pmatrix} = \begin{pmatrix} 0 & 0 & 0 & 1 \\ 0 & 1 & 0 & -1 \\ 0 & 0 & 1 & -1 \\ 1 & -1 & 1 & 1 \end{pmatrix} \cdot \begin{pmatrix} \pi_{2q} \\ \pi_{12} \\ \pi_{23} \\ \pi_{23} \end{pmatrix} \]  

(Eq. 5)

For \([1,6-^{13}C_2] \)glucose perfusion, the fitted time courses were as follows: Glu4Tot (Tot-total), Glu4s, Glu4d34, Glu3Tot, Glu3d, Glu2Tot, Glu2s, Lac3Tot, and acylmethylene (\(-\text{CH}_2\)) resonance, for a total of nine curves. The following free fluxes were determined: melanoma TCA cycle \( F_{\text{TCA}} \); pyruvate carboxylase \( F_{\text{PC}} \); exchange between glutamate and \( \alpha \)-ketoglutarate \( F_X \); anaplerotic exchange and net flux at the level of succinyl-CoA; \( F_{\text{ANA}} \); mitochondrial malic enzyme (ME2+ME3) activity; \( \text{de novo} \) fatty acid production, glutaminolysis, aspartate efflux; and three other parameters, Michaelis-Menten lactate transport \( V_{\text{max}}^{\text{LAC}} \), and \( K_{m}^{\text{LAC}} \), and total cellular lactate concentration. Other fluxes and parameters, including the Warburg effect value, were calculated based on flux balance analysis, including non-oxidative glycolytic flux CMRlac and others. Solving a system of differential equations (using the Runge-Kutta 4th order procedure for stiff systems) in terms of bonded cumomer variables yields time courses for all possible fine structure \( ^{13}C \) multiplets of glutamate, glutamine, and aspartate.
The classical PPP consists of two branches, irreversible oxidative and reversible non-oxidative parts. The oxidative branch represents the net PPP flux and produces NADPH, which can be used in many reductive synthetic pathways, including de novo lipogenesis. The PPP reactions are summarized in the Fig. 2.

To describe the pentose phosphate pathway, we used the steady-state bonded cumomer flux model variant (which results in a system of non-linear algebraic equations for the $\pi_{(i)}$ cumomer fraction; see the equation for $\pi_{(i)}$ instead of the set of differential equations) to fit $^{13}$C NMR lactate data acquired after an 8-h incubation of DB-1 cells with $[1,2-^{13}$C$_2]$glucose tracer. The fitted steady-state data were Lac3Tot, Lac2Tot, Lac3s, Lac3d23, Lac2s, and Lac2d23 multiplets. The relations between lactate multiplets and bonded cumomers are presented elsewhere (22).

Non-oxidative reactions are assumed to follow carbon atom redistributions (based on the ping-pong chemical mechanism) depicted in the following atom distribution matrices (ADM) (22, 35). Particularly for the transketolase 1 (TK1) reaction, $X5P \leftrightarrow G3P + S7P$, where xylulose 5-phosphate ($X5P$) and ribose 5-phosphate ($R5P$) forms glyceraldehyde 3-phosphate ($G3P$) and sedoheptulose 7-phosphate ($S7P$), the corresponding ADM is shown in Matrix 1.

\[
\begin{array}{cccccccc}
1 & 2 & 3 & 4 & 1 & 2 & 3 & 4 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\
\end{array}
\]

In this matrix the symbol “1” reflects a particular carbon transfer from substrates (the upper level of the matrix indicates carbon atoms of $X5P$ and the lower level indicates carbon atoms of $R5P$) to products (numbers in the 1st row reflect carbon atoms of $G3P$ from 1 to 3 and $S7P$ from 1 to 7), and zero reflects no channeling between particular atoms in the metabolites. For the transaldolase reaction, $S7P \leftrightarrow G3P + E4P$, where sedoheptulose 7-phosphate ($S7P$) forms glyceraldehyde 3-phosphate ($G3P$) and erythrose 4-phosphate ($E4P$), the corresponding ADM is shown in Matrix 2.

\[
\begin{array}{cccccccc}
1 & 2 & 3 & 4 & 1 & 2 & 3 & 4 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\
\end{array}
\]

We accounted for the reversibility of every reaction by considering forward and reverse reaction steps through the non-oxidative PPP branch with a different extent of reaction reversibility for every reaction above. The system will still result in constant oxidative PPP net flux for every reaction. Overall, these matrices show that label at a particular 1st position of pentose 5-phosphate in the oxidative branch will result in the labeling of all carbon atoms in downstream metabolites such as pyruvate and lactate.

Metabolic and transport parameters as well as metabolite pool sizes used here are published elsewhere (36). For other parameters such as concentrations of the TCA cycle intermediates not reported in this reference, pool sizes were assumed to be in the range 0.1–0.2 mM. The total citrate pool size was assumed to be 0.5 mM with 20% in the mitochondria and 80% in cytosol. Pyruvate was assumed to be 20% in mitochondria. Sensitivity analysis (not shown here) verified that the chosen TCA cycle intermediate pool sizes did not affect extracted fluxes within 5% (even when modified by factors of 5 both above and below the assumed values).

Minimization of sum-of-squared residuals between simulated and experimental data was performed using Broyden-Fletcher-Goldfarb-Shanno, Levenberg-Marquardt, or Simplex algorithms. Proper mean square convergence was confirmed by verifying that goodness-of-fit values were close to expected theoretical values. The errors of the calculated parameters were estimated using Monte Carlo simulations with experimental noise levels (37). All numerical procedures were carried out in Matlab (Mathworks, Natick, MA).

In this study, LC-MS data have been utilized to compare fluxes under different conditions (e.g. hyperglycemia versus euglycemia) without detailed flux analysis (because no statistically significant change was observed between the different medium glucose conditions).
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![Diagram of Bioreactor Design and NMR Spectra of DB-1 Melanoma Cells](image)

**Results**

$^{31}P$ NMR—A typical $^{31}P$ spectrum is shown in Fig. 1B. The $\beta$-NTP resonance at $-16.5$ ppm was used to estimate the cell concentration. Other prominent resonances in the spectrum are $\alpha$- and $\gamma$-NTP, diphosphodiesters, glycerophosphocholine, glycerophosphoethanolamine, inorganic phosphate, phosphocholine, phosphoethanolamine, and diphosphodiesters, including NADP(H). A broad hump emanating from immobilized phosphates from lipids dominates the central region of the spectrum. This broad peak can be removed by off-resonance saturation that is mediated by spin diffusion through the phospholipid semi-immobilized matrix, but this might modify resonance intensities of metabolites that are partially or transiently bound to membranes (38). We therefore have avoided off-resonance saturation. Other methods of removing the “hump” include fitting it with a spline function and subtracting the base line or subtracting the first spectrum and monitoring differences in peak intensities over time (39). The spectrum in Fig. 1B indicates the energy level of the tumor cells (measured by the intensity of the $\beta$-NTP peak). The level was measured before and after the time course of the $^{13}C$ NMR experiment and was unchanged. Usually, the bioenergetic status of the cells is measured by the $\beta$-NTP/P$_{i}$ ratio; however, in a bioreactor the P$_{i}$ resonance is dominated by the P$_{i}$ in the perfusate. The latter peak could be removed by applying strong gradients that dephase resonances of flowing metabolites (38, 40), but we have not applied this method here because it could lead to errors if removal of mobile resonances was incomplete.

**Metabolic Fate of [1,6-$^{13}C_2$]Glucose in DB-1 Cells**—Fig. 2 shows the entire metabolic network that is included in the current analysis (see under “Experimental Procedures” for details). $^{13}C$ labeling results for DB-1 melanoma cells grown at a constant glucose level of 26 mM are shown in Figs. 1C, 3, and 4. Fig. 1C displays a $^{13}C$ NMR spectrum that was obtained during the final 15 min of the experiment. Resonances were detected for C3 of lactate, C2, C3, and C4 of glutamate, C2 and C3 of aspartate, and fatty acyl carbons. With the glucose concentration clamped at 26 mM, lactate labeled at C3 (20.9 ppm) appeared in the spectrum within the first few minutes (Fig. 4A). The level of this resonance increased throughout the experiment but did not reach saturation. The time course of the intensity of this resonance is the combined result of the biological rate of lactate formation and the rate of washout of lactate due to the continuous addition of fresh medium and withdrawal of spent medium from the perfusion system. Fifteen minutes after introduction of labeled glucose, $^{13}C$ incorporation was detected in C4 of glutamate (34.3 ppm); the full time course for labeling results for this resonance is shown in Fig. 3. The concentration increased in a linear manner for approximately 1 h. Subsequently, satellites were observed in this resonance indicating the presence of $^{13}C$-$^{13}C$ coupling between glutamate-C4 and glutamate-C3 (5). The rate of labeling of glutamate exclusively on C4 gradually slowed and reached close to steady state after ~5 h. The level of glutamate labeled at both C3 and C4 continued to increase steadily throughout the experiment. Fig. 3, A and B, shows the labeling results for the glutamate-C3 resonance at 28.8 ppm. Over the 1st h, essentially no $^{13}C$ labeling was detected in glutamate-C3. Subsequently, $^{13}C$ labeling of glutamate-C3 increased linearly but did not reach saturation. Approximately two-thirds of the C3 glutamate carbon was coupled (predominantly to glutamate-C4). For glutamate-C2, label also first appeared approximately 1 h after the beginning of the experiment (Fig. 3, A and B). However, unlike glutamate-C3, only a singlet was observed for the first 3 h. Subsequently, a doublet appeared due to the presence of glutamate-C2,3 labeling. Such glutamate would be produced after carbon 1 was removed from glutamate-C3,4 during the third turn of the TCA cycle (5).
Labeling was also observed in long chain fatty acids at ~30 ppm, indicating that DB-1 melanoma cells produce lipid de novo from glucose (see Fig. 4B). This result is consistent with results for other cancer cell lines grown in culture (19). Small amounts of labeling were observed in aspartate-C2 and C3. No label was observed for any carbon of alanine, probably due to low total concentrations of alanine; this result is unusual because most cancer cell lines produce 13C-labeled alanine from glucose (19, 20). Resonances were observed in regions associated with the glycerol backbone of phospholipids in triglycerides at ~65 ppm. This region also contains resonances associated with glycolytic intermediates.

Quantifying Bionetwork Fluxes—Our biochemical network model was used to fit 13C NMR DB-1 melanoma data acquired during continuous perfusion of cells with [1,6-13C2] glucose in the bioreactor depicted in Fig. 1A. The experimental data from isotopomer time courses was converted to 13C-metabolite concentration curves (Figs. 3 and 4). Furthermore, to minimize the bias introduced by using multiplets with low signal to noise ratio, only those multiplets with end point concentrations greater than 0.5 mM were used in the fitting procedure. The remaining experimental multiplets were used as reference points for visually examining the fit predicted for low intensity curves to experimental data but not actually calculating the goodness of fit of these metabolite resonances, whose noise levels were expected to be high (not shown here). To quantify fluxes in central metabolism, we combined four types of measurements as follows: three sets of time courses of 1) glutamate isotopomers (multiplets); 2) total lactate-C3 labeling; 3) acyl groups of fatty acid resonances determined by NMR in situ; and 4) steady-state 13C NMR lactate isotopomers. The fitted time courses were as follows: Glu4Tot (where Tot is total), Glu4s, Glu4d34, Glu3Tot, Glu3d, Glu2Tot, Glu2s, Lac3Tot, and acyl methylene (–(CH2)n–) resonance, yielding a total of nine curves. Estimation of the model-free parameters was achieved by fitting labeled patterns of glutamate, lactate, and fatty acid acyl methylene groups. The extracted flux values are presented in Table 1 and in the flux map (see Fig. 5). Flux confidence intervals were calculated by Monte-Carlo simulations (see Fig. 6A, only four fluxes are shown) and are summarized in Table 1.

Quantifying ATP Production Routes—The predicted fluxes were used to calculate contributions of glycolysis and oxidative phosphorylation to ATP production. We assumed that the oxidative phosphorylation phosphate to oxygen ratio equals 2.5 for NADH and 1.5 for FADH2 oxidation. To assess the contribution of glucose, fatty acids (FA) and glucogenic amino acids, and glutamine to generating reducing power, we calculated the fluxes for each of these metabolites. For each of the above substrates, we calculated the NADH/FADH2 reaction flux based on the different labeling patterns that we were able to track both...
from the labeled and unlabeled carbon atoms. The combined cytosolic ATP production flux was 150 mmol/liter-cell/h, and the mitochondrial compartment produced 154 mmol/liter-cell/h due to oxidative phosphorylation. The results are presented in Table 1.

**Pentose Phosphate Pathway (PPP)—** We fitted $^{13}$C NMR steady-state lactate multiplets acquired after an 8-h incubation of DB-1 cells with [1,2-$^{13}$C$_2$]glucose tracer. The experimental $^{13}$C NMR spectra and the results of fitting are shown in the Fig. 7. The extracted fluxes relative to glycolytic flux are presented in Table 1. Flux standard deviations were calculated by Monte-Carlo calculations. Flux through the oxidative PPP was estimated to be 3.6% of the glycolytic rate. Given the extensive experimental data that have been obtained for the lactate $^{13}$C NMR multiplets, we were also able to estimate non-oxidative exchange fluxes (the smaller rates of each reversible reaction) of the classical PPP (Fig. 2), which are presented in Table 1. Transketolase 1 and transaldolase exchange fluxes in the non-Ox PPP branch were consequently relatively high at 21 and 44%, respectively, and transketolase 2 activity was close to zero. The flux standard deviations (Table 1) demonstrate that all fluxes through the PPP were resolved but with relatively poor resolution (when using only $^{13}$C NMR isotopomers of lactate as a reporter molecule). NADPH production flux via the oxidative PPP pathway was estimated to be 6.6 mmol/liter-cell/h. At the end of this experiment, we measured glucose and lactate concentrations in the flask experiments. Glucose concentrations under euglycemic conditions were unchanged within 30%.

**Quantifying Glutaminolysis Flux—** Our recent focus has turned to refinement of the details of glutaminolysis because of its critical role in the metabolism of many malignancies. For example, one glutamine molecule could produce up to 22.5 ATP molecules. Therefore, even a small amount of glutaminolysis can have a major effect on the bioenergetics of cancer cells. In DB-1 melanoma under hyperglycemic conditions, glutamine uptake flux ranged up to 50% of the TCA cycle flux.
Under euglycemic conditions, glutamine uptake doubled reaching approximately the same level as TCA cycle flux. Glutamine uptake can lead to protein synthesis, glutathione synthesis, or nucleotide synthesis as well as to ATP production by the glutaminolysis pathway (see below) or to cytosolic reductive carboxylation driven by glutaminase 1 (GLS; it is considered to be present both in mitochondria and cytosol) or other glutamine-glutamate conversion enzymes in cytosol and isocitrate dehydrogenase 1 (IDH1) (41). It was, therefore, important to refine the metabolic analysis of DB-1 melanoma to accurately assess the contributions of the glutaminolysis pathway in these tumor cells (see Fig. 2). To quantify the glutaminolysis contribution to energy production, we modified the bionetwork by simultaneously allowing net glutamine uptake.

**TABLE 1**

Extracted fluxes using bonded cumomer modeling

| Reaction | Flux*<sub>a</sub> (mmol/liter-cell/h) |
|----------|----------------------------------|
| Pentose phosphate pathway (oxidative branch), net PPP F_{PPPox} | 3.3 ± 72% |
| Transketolase 1 (exchange) relative to glycolysis, TK1 | 21 ± 56% |
| Transketolase 2 (exchange) relative to glycolysis, TK2 | 0.5 ± 90% |
| Transaldolase (exchange) relative to glycolysis, TA | 44 ± 64% |
| Glucogenic amino acids contribution to pyruvate, F_{FGAA} | 5.0 ± 20% |
| TCA cycle rate, F_{TCA} (isocitrate → α-Kg) | 10.9 ± 8% |
| Exchange, Mal-Asp shuttle, F_{mal} (reversible) | 11.3 ± 19% |
| Pyruvate carboxylase flux, F_{pc} | 0.6 ± 51% |
| Anaplerotic exchange flux at succinyl-CoA, F_{anap} | 3.0 ± 15% |
| Net anaplerotic flux to the TCA cycle, F_{anap} | 4.3 ± 15% |
| Flux of ketogenic AA and FA to acetyl-CoA formation, F_{anap} | 0.1 ± 19% |
| Glutaminolysis flux, F_{gls} | 0.05 ± 60% |
| Pyruvate dehydrogenase flux, F_{pdh} | 10.8 ± 16% |
| Citrate synthase flux, F_{cs} | 11.5 ± 14% |
| Net fumarase activity, F_{fum} | 15.3 ± 16% |
| Fatty acid synthesis, F_{fas} | 0.59 ± 15% |
| Mitochondrial pyruvate transport, F_{mpc} | 7.1 ± 23% |
| Malic enzyme (mitochondrial), ME2 + ME3, F_{mem} | 4.3 ± 12% |
| Malic enzyme (cytosolic), ME1, F_{me} | 0.6 ± 15% |
| Aspartate efflux, F_{as} | 0.06 ± 45% |
| Production flux of ATP, glycolytic, Warburg, F_{ATP,g} | 150 ± 13% |
| Production flux of ATP, mitochondrial, oxphos, F_{ATP,o} | 154 ± 12% |
| Warburg parameter (net LDH/MPC flux ratio) | 21 ± 28% (unit-less) |
| Combined glycolysis and PPP flux from G6P to pyruvate | 152.1 ± 12% |
| Glutamine consumption flux (for protein, nucleotides, amino acids, glutathione, etc.) | 4.9 ± 10%<sup>b</sup> |
| Net LDH activity, Pyr → Lac, F_{LDH} | 150 ± 12% |
| Lactate transport, V_{maxlact} | 395 ± 12% |
| Lactate transport, Michaelis-Menten parameter, K_{m} | 3.5 mM ± 70% |

<sup>a</sup> Calculated fluxes are expressed in mmol/liter-cell/h with relative standard deviations in percent for pathways shown in Fig. 2. Warburg parameter is the ratio of the net flux through lactate dehydrogenase (LDH) to the mitochondrial pyruvate carrier influx (MPC).

<sup>b</sup> Standard deviation was estimated based on experimental measurements of glutamine consumption.

FIGURE 5. Metabolic flux map and energy metabolism with predicted oxygen consumption rate MRO2 in the human melanoma DB-1 bionetwork at hyperglycemia and normoxia. Numbers indicate net fluxes in mmol/liter-cell/h. See Table 1 for definitions and values of the derived fluxes. Reversibility for SDH, Fumarase, and MDH was taken into account as 10% of the direct flux. Varying the extent of reversibility in the range 5–30% does not change extracted fluxes within several percent (results not shown).
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There are two pathways for pyruvate-malate cycling. One pathway involves conversion of malate via mitochondrial malic enzyme (ME2) to pyruvate; it then returns to oxaloacetate via pyruvate carboxylase or it can go from pyruvate to acetyl-CoA leading to citrate. The second pyruvate cycling pathway goes from malate in the mitochondria through a transporter into the cytosol and from there to cytosolic pyruvate via cytosolic malic enzyme (ME1). The cytosolic pyruvate can return to the mitochondria through the mitochondrial pyruvate carrier or it can be converted to alanine via alanine-aminotransferase (GPT1) or to lactate via lactate dehydrogenase. The BC approach can distinguish between these pyruvate-malate pathways or their combinations under real conditions (Fig. 2).

“Classical” glutaminolysis includes uptake of glutamine by the cell, transport to mitochondria, conversion to glutamate through mitochondrial glutaminase 1 and 2 (GLS and GLS2), entry into the TCA cycle to α-ketoglutarate via glutamate dehydrogenases 1 and 2 (GLUD1 and GLUD2) or via mitochondrial aminotransferases (GPT2 and GOT2), traversing to oxaloacetate, and exiting as pyruvate or aspartate. Alternatively, glutaminolysis could partially occur in the cytosol by producing glutamate or α-ketoglutarate from glutamine and eventually by entry of those metabolites into the TCA cycle as α-ketoglutarate. Further flux in the TCA cycle then moves to malate where malate-pyruvate cycling occurs (Fig. 2). There are two pathways for pyruvate-malate cycling. One pathway involves conversion of malate via mitochondrial malic enzyme (ME2) to pyruvate; it then returns to oxaloacetate via pyruvate carboxylase or it can go from pyruvate to acetyl-CoA leading to citrate. The second pyruvate cycling pathway goes from malate in the mitochondria through a transporter into the cytosol and from there to cytosolic pyruvate via cytosolic malic enzyme (ME1). The cytosolic pyruvate can return to the mitochondria through the mitochondrial pyruvate carrier or it can be converted to alanine via alanine-aminotransferase (GPT1) or to lactate via lactate dehydrogenase. The BC approach can distinguish between these pyruvate-malate pathways or their combinations under real conditions (Fig. 2).

We continue the analysis of data of DB-1 melanoma cells treated with [1,6-13C2]glucose as the tracer. Dynamic analysis indicates that the curves resulting from fitting of the 4d34-labeled glutamate multiplets, particularly the glutamate 3,4-doublet, Glu4d34, resulting from simultaneous labeling of C3 and C4 (during the second TCA cycle turn) and the singlet resulting from glutamate labeling exclusively at C4 (during the first turn of the TCA cycle), are very sensitive to flux through the glutaminolysis pathway even when only natural abundance medium-derived glutamine has been used. See for example Fig. 6B, where the separation and shape of the glutamate C4 total and C4 singlet and 3,4-doublet, Glu4d34, curves...
The mass isotopomer distribution of several TCA flux (see Table 1). As validation of the inferred/extracted fluxes, we measured the MR\textsubscript{Glc} with polarographic oxygen probes during the perfusion experiment. The fluxes obtained by fitting of dynamic \textsuperscript{13}C data produced good agreement between the experimentally observed MR\textsubscript{Glc}, 33 mmol/liter-cell/h, and the calculated MR\textsubscript{Glc}, 32 mmol/liter-cell/h (see Table 2). The experimental total glutamate concentration was measured at the end of the bioreactor experiment by cell extraction and LC-MS analysis.

We then evaluated the contribution of glutaminolysis flux to overall energy production of DB-1 melanoma cells. We found that for this tumor line, glutaminolysis flux through the truncated TCA cycle and via pyruvate-malate cycling by malic enzyme ME2 is essentially zero. This conclusion resulted directly from the ability of the model to detect glutaminolysis flux by fitting of data on the glutamate multiplets (see above). To further confirm this conclusion, we forced the model to accommodate glutaminolysis flux between 5 and 20% of the TCA cycle flux. Invariably, this grossly deteriorated the goodness-of-fit of the isotope kinetic data to the model (results not shown).

Even though glutaminolysis does not contribute to the mitochondrial ATP production, there is still substantial mitochondrial malic enzyme flux (combined activities of NADH producing ME2 and NADPH producing ME3), which is equal to 40% of the TCA flux (see Table 1).

**Effect of Hyperglycemia/Euglycemia on Glutamine and Fatty Acid Metabolism**—The mass isotopomer distribution of several TCA cycle metabolites from \textsuperscript{[\textsuperscript{13}C\textsubscript{5}, \textsuperscript{15}N\textsubscript{2}]}glutamine labeling experiments is shown in Fig. 8 at different glucose concentrations. With this tracer, increased glutaminolysis flux resulted in increased abundance of the M + 4 mass isotopomer of citrate, succinate, and malate (41), whereas reductive carboxylation resulted in increased M + 3 for malate, M + 4 for succinate, and M + 5 for citrate. Reductive carboxylation and increased glucose contribution to the fatty acid production also disturbs the acetyl-CoA mass isotopomer distribution. Fig. 8 clearly shows that there are no significant differences in labeling patterns of all four metabolites. Consequently, based on bioreactor data, we can exclude any major change in glutaminolysis flux as well as for fatty acid production under hyperglycemic versus euglycemic conditions.

**Quantifying de Novo Lipogenesis Flux in DB-1 Cells**—During culturing of DB-1 cells with \textsuperscript{[1,6-\textsuperscript{13}C\textsubscript{2}]}glucose, progressive labeling of fatty acids was observed for multiple resonances (see Figs. 1C and 4B). FA synthesis occurs first by cytosolic citrate production from glucose and through reductive carboxylation in the cytosol from glutamine through isocitrate dehydrogenase (IDH1 isoform). By ATP citrate lyase, cytosolic citrate is converted to oxaloacetate and cytosolic acetyl-CoA, which is converted further to malonyl-CoA by the rate-limiting acetyl-CoA carboxylase (ACC-B) and through fatty acid synthases to elongated precursors via sequential addition of acetyl-CoA, eventually leading to palmitate production. Palmitate is further elongated or desaturated to produce numerous other FAs. To address de novo FA flux production, we utilized the BC model with cytosolic FA and their precursors as well as membrane fatty acids as the second FA pool. We calculated FA content based on the total FA content of ~20% of cell dry weight of 200 mg/10\textsuperscript{9} cells (5  \times  10\textsuperscript{8} cells = 1 ml). Total FA content referenced to palmitate is at the level of several tens of mmol/liter-cell. We iteratively varied the ratio of cytosolic to membrane FA and found the best fit at ~10 times more FA in the cytosolic membrane compartment (Fig. 4B). The de novo flux of conversion of glucose to FA was 0.59 mmol/liter-cell/h, and NADPH consumption flux in FA biosynthesis was 8.4 mmol/liter-cell/h.

**Model Validation**—As validation of the inferred/extracted fluxes, we measured the MR\textsubscript{Glc} with polarographic oxygen probes during the perfusion experiment. The fluxes obtained by fitting of dynamic \textsuperscript{13}C data produced good agreement between the experimentally observed MR\textsubscript{Glc}, 33 mmol/liter-cell/h, and the calculated MR\textsubscript{Glc}, 32 mmol/liter-cell/h (see Table 2). The experimental total glutamate concentration was 8.6 mmol/liter-cell/h. Assuming the glutamate concentration to be a fitted
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**TABLE 3**

| Glutamine | Glucose | Glucose flux | Glucose flux |
|-----------|---------|--------------|--------------|
| Euglycemic (6.4 mM) | 0.064 | 0.86 | |
| Hyperglycemic (23.2 mM) | 0.035 | 0.79 | |

Experimental glucose and glutamine consumption fluxes under euglycemic (6.4 mM) and hyperglycemic conditions (23.2 mM in medium).

Glutamine = 3.0 mM at low glucose and 3.4 mM at high glucose. F_Glc, glucose consumption flux (mmol/10^6 cells/h) and F_Gln, glutamine consumption (mmol/10^6 cells/h). Flux ratios were F_Glc/F_Gln = 22.4 at high glucose and F_Glc/F_Gln = 13.5 at euglycemia.

Under euglycemic conditions, glutamine influx approximately doubles, whereas glucose consumption remains essentially unchanged. Table 4 shows the glucose and lactate concentrations at the end of experiments at different glycemic conditions conducted in the flask experiments for mass spectrometric measurements. Glucose concentrations were unchanged within 30% under euglycemic conditions and within 5% under hyperglycemic conditions. Finally, we were able to generate a flux map (Fig. 5). This figure shows the extracted metabolic fluxes with the predicted MR_O2. Numbers indicate fluxes in mmol/liter-cell/h. See Table 1 for definitions and values of the derived fluxes.

**Discussion**

Measuring the relative glycolytic and oxidative capacity of specific tumors plays an important role in development of strategies for treating neoplasms and for monitoring tumor response to specific therapeutic agents. For this reason, development of a non-invasive method for quantifying the fraction of ATP production by tumors that occurs by glycolytic and mitochondrial metabolism would be useful in the management of cancer and tailor-fitting of therapeutic regimens to the needs of individual patients. In addition, various other pathways of tumor metabolism play critical roles in tumor proliferation by providing additional energy sources and essential precursors for cell replication. Our goal was to develop a method for quantitating tumor intermediary metabolism in cells and with some extension in animal tumor models and human cancer patients.

One of our key objectives was to validate the bonded cumomer modeling technique as a method for measuring fluxes through key pathways of tumor energy production. This is best accomplished by comparing model predicted parameters with independently measured experimental parameters as has been demonstrated previously by Chatham et al. (6) for the cardiac TCA cycle model of Chance et al. (5) and Jeffrey et al. (7) for the isotopomer model of the heart. Table 2 compares model-predicted and experimentally measured parameters. There is excellent agreement between the predicted MR_O2 (32 mmol/liter-cell/h) and the experimentally measured rate of oxygen consumption (36 mmol/liter-cell/h). The experimental and predicted rates of glucose uptake agree within 7%. The experimental and predicted pyruvate to lactate fluxes, 149 and 150, respectively, agree within ~1% and glutamate pool sizes within 7%. Overall, this excellent agreement between model-predicted and experimental data validates the BC model.

ATP production fluxes were calculated assuming a phosphate to oxygen ratio of 2.5 for NADH and 1.5 for FADH2 (42). Under hyperglycemic conditions, the average ATP production rates were MR_ATP (mitochondrial) = 154 mmol/liter-cell/h and MR_ATP (glycolytic) = 150 mmol/liter-cell/h which indicates that 51% of the energy was produced by oxidative phosphorylation and 49% was from glycolysis. Based on the similarity of isotopomer labeling patterns under hyperglycemic and euglycemic conditions (Fig. 8), we expect no significant change in glycolytic and oxidative metabolism under these conditions. High fluxes of pyruvate mitochondrial transport (~70% of the TCA cycle activity) and mitochondrial ATP production are promoted by the melanoma-associated BRAF mutation. Recent work (43) has demonstrated that BRAF V600E mutated melanoma cells exhibit increased pyruvate entry through the activated PDH complex. PDH flux (~106% of the TCA flux) is regulated by inhibitory phosphorylation mediated by PDH kinases (PDK) and PDH phosphatases (PDP). The oncogene BRAF V600E causes concerted activation of the PDH complex by down-regulation of PDK1 expression and up-regulation of PDP2 thereby promoting oxidative pyruvate utilization and leading to increased cellular respiration.

There has been considerable interest in glutamine metabolism as a key source of tumor energy production linked to c-Myc expression (21, 44 – 47). Our model of tumor metabolism (Fig. 2) contains a specific glutamine transporter on the cell membrane. The classical model of glutaminolysis includes an additional specific glutamine transporter on the mitochondrial membrane that leads to mitochondrial glutaminase (GLS2 or GLS), which converts glutamine to glutamate, which is then converted to α-ketoglutarate by enzymes such as glutamate dehydrogenase (GLUD1 and -2). Two additional transaminases can carry out the glutamate-α-ketoglutarate exchange, aspartate aminotransferase, and alanine aminotransferase. The α-ketoglutarate then produces reducing equivalents via the oxidative branch of the TCA cycle leading to ATP production by oxidative phosphorylation. Various pathways, including the malate-aspartate shuttle, pyruvate carboxylase, malic enzyme, and phosphoenolpyruvate carboxykinase (PEPCK1 and -2) all
Conclude that glutamine was utilized as an anaplerotic substrate for making this crucial distinction. Our labeled LC-MS data for production rate. Bonded cumomer analysis provides a method of these two substrates. The data fitting clearly indicated that the net flux from glutamine being metabolized via the TCA cycle was very small.

We also conducted a sensitivity analysis to show that using \([1,6-^{13}C_2]\)glucose tracer alone suffices to reliably calculate glutaminolysis flux in malignant cells and tissues. Fig. 6B shows that the glutamate C4 doublets due to C3-C4 coupling (Glu4d34) are sensitive to net flux of unlabeled glutamine entering the TCA cycle; even a 10% difference in flux produces a clearly detectable shift in these curves. Fitting analysis indicated that there was little net influx from glutamine into the TCA cycle. Note also that this conclusion can also be reached from \(^{13}C\) NMR data without the use of labeled glutamine.

By contrast, some investigators utilizing mass spectrometry have equated glutamine uptake minus glutamate release with net glutaminolysis flux (48). In this case, the net influx of glutamine is \(\sim 5\) mmol/liter-cell/h, which, if it were equated with glutaminolysis flux, would have been interpreted as 113 mmol/liter-cell/h of ATP per h or about 75% of the glycolytic production of ATP. Under euglycemic conditions, which produce about twice the level of glutamine uptake, such an analysis would have attributed about 50% higher ATP production to glutaminolysis than to glycolysis, and most of the tumor energy production would have been attributed to glutaminolysis. This is incorrect in our case; in reality the net glutaminolysis flux under hyperglycemic conditions is \(\sim 0.1\) mmol/liter-cell/h or \(\sim 2\) mmol of ATP/liter-cell/h or \(\sim 1\)% of the glycolytic ATP production rate. Bonded cumomer analysis provides a method for making this crucial distinction. Our labeled LC-MS data for the TCA cycle intermediates with \([U-^{13}C_2,U-^{15}N_2]\) glutamine confirmed that no glutaminolysis occurred under euglycemic conditions versus hyperglycemia (Fig. 8). The absence of change in glutaminolysis or reductive carboxylation flux is evident from similar mass isotopomer distributions of several key metabolites. Our finding of nearly zero glutaminolysis flux contradicts Scott et al. (49), who, on the basis of glutamine uptake and label appearance in some TCA cycle metabolites, concluded that glutamine was utilized as an anaplerotic substrate of several human melanoma lines. However, these authors did not conduct a rigorous metabolic modeling analysis to measure flux through the TCA cycle. Consequently, they were unable to distinguish between actual flux through the cycle via glutaminolysis and exchange labeling between \(\alpha\)-ketoglutarate and glutamate without any net flux. In addition, glutamine could be used for many other metabolic processes besides energy production, e.g. protein, nucleotide, and hexosamine synthesis.

If the net uptake of glutamine is 5 mmol/liter-cell/h with negligible contribution to energy production, glutamine must be contributing primarily to anabolic processes such as amino acid, nucleotide, and protein production as well as to \(de novo\) lipogenesis. Although glutamine-derived carbons have been unlabeled, we have monitored lipogenesis by \(^{13}C\) NMR detecting methylene carbon acyl groups coming from glutamine (Figs. 1C and 4B). Table 1 indicates that \(de novo\) lipogenesis corresponds to \(\sim 6\)% of the TCA cycle flux under hyperglycemic conditions. However, this does not include possible direct contributions from unlabeled glutamine (i.e. through cytosolic reductive carboxylation by isocitrate dehydrogenase IDH1 or mitochondrial IDH2) and potentially from the cytosolic acetyl-CoA synthetase 2 (ACSS2) reaction (50).

This modest flux for \(de novo\) fatty acid synthesis requires a large contribution of NADPH cofactor production (\(\sim 8.3\) mmol/liter-cell/h). NADPH could be produced by the oxidative branch of the PPP, malic enzyme, or isocitrate dehydrogenase activities or by serine-glycine one-carbon metabolism (folate metabolism). Our calculated malic enzyme activities in mitochondria (even if all this activity is represented by ME3) would account for 4.3 mmol/liter-cell/h, and cytosolic NADPH flux (ME1, 0.6 mmol/liter-cell/h) would not account for the required high level of NADPH flux. A recent elegant study (51) suggested that in some cancers folate metabolism is the largest source of NADPH. Based on these data (52), we estimated that the serine-glycine one-carbon metabolism pathway cannot account for the required NADPH consumption. In the light of our \(^{13}C\) NMR lactate labeling data, the oxidative branch of the PPP is the main source of NADPH in these cancer cells (6.6 mmol/liter-cell/h). This constitutes \(\sim 80\)% in agreement with the NADPH consumption rate during fatty acid biosynthesis. Thus, the flux through the oxidative PPP appears to be \(\sim 80\)% sufficient to supply all of the NADPH required for fatty acid production and is the main supplier of reducing equivalents in DB-1 melanoma. A 20% difference in the rates of NADPH production and consumption indicates that alternative sources of NADPH+ reduction, considered above, need to be taken into account. We also found that mitochondrial pyruvate-malate cycling was very high (4.3 mmol/liter-cell/h). This could also contribute to defense against reactive oxygen species by producing sufficient NADPH via ME3.

In summary, we have validated a novel method, bonded cumomer analysis, for determining fluxes through specific pathways of tumor energy metabolism. This method can be applied to studies of perfused tumor cells in bioreactors or, with some modifications, to \(in vivo\) studies for non-invasive investigation of the metabolism of tumors in animals or humans. The transition from isolated cells to \(in vivo\) rodent models and humans is not simple and seamless, but it is feasible. The interested reader experienced in metabolic modeling might pursue similar objectives using the methods outlined in this study together with mathematical methods and metabolic models referenced herein (53).
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Author Contributions—The study was conceived and coordinated by J. D. G., A. M., and A. A. S.; A. M., S. C. L., L. G., D. S. N., and J. C. R. designed and performed the experiments and collected the data; A. A. S., A. M., S. C. L., I. A. B., and J. D. G. designed, analyzed, and interpreted the experiments; A. A. S. performed computational metabolic analysis; P. G. H. and D. B. L. provided technical assistance and contributed to the preparation of the manuscript; J. D. G. provided conceptual advice; A. A. S., A. M., S. C. L., and J. C. R. designed and performed the experiments and collected the data; A. A. S. performed computational metabolic analysis; P. G. H. and D. B. L. provided technical assistance and contributed to the preparation of the manuscript; J. D. G. wrote and edited the manuscript with the assistance of all authors.

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