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Gustafsson, J., Roshanzamir, F., Hagnestål, A. et al (2024). Metabolic collaboration between cells in the tumor microenvironment has a negligible effect on tumor growth. *Innovation*, 5(2). <http://dx.doi.org/10.1016/j.xinn.2024.100583>

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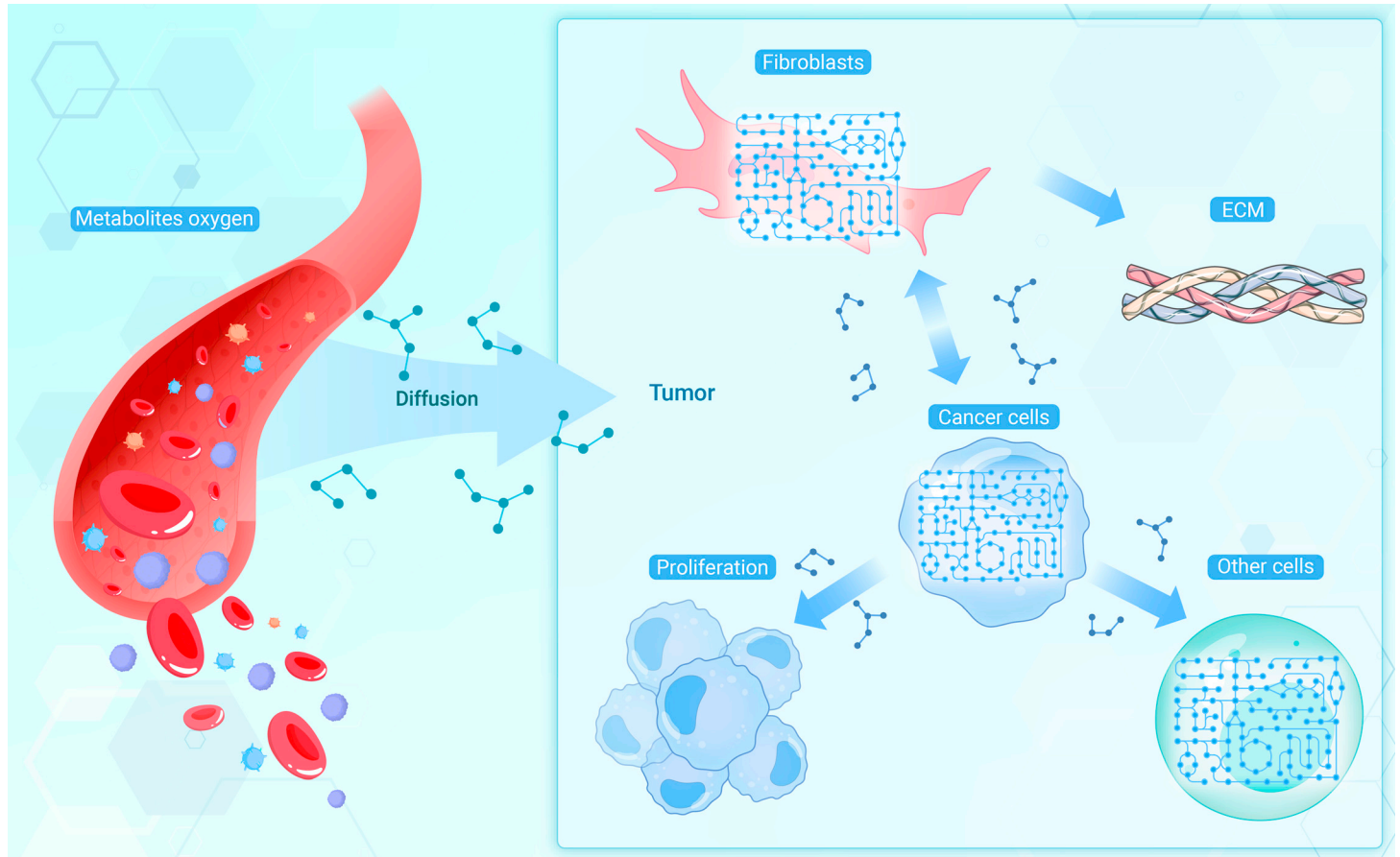
Johan Gustafsson,¹ Fariba Roshanzamir,¹ Anders Hagnestål,² Sagar M. Patel,³ Oseeyi I. Daudu,³ Donald F. Becker,³ Jonathan L. Robinson,^{1,4} and Jens Nielsen^{1,4,*}

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Received: April 28, 2023; Accepted: January 24, 2024; Published Online: January 30, 2024; <https://doi.org/10.1016/j.xinn.2024.100583>

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GRAPHICAL ABSTRACT



PUBLIC SUMMARY

- Simplified application of enzyme usage constraints to genome-scale metabolic models with the software GECKO Light.
- Estimation of maximum metabolite influx into tumors at varying levels of hypoxia.
- Simulations of tumor metabolism explain “glutamine addiction” in cancers.
- Metabolic collaboration between cell types in the tumor microenvironment does not increase tumor growth.



Metabolic collaboration between cells in the tumor microenvironment has a negligible effect on tumor growth

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The tumor microenvironment is composed of a complex mixture of different cell types interacting under conditions of nutrient deprivation, but the metabolism therein is not fully understood due to difficulties in measuring metabolic fluxes and exchange of metabolites between different cell types *in vivo*. Genome-scale metabolic modeling enables estimation of such exchange fluxes as well as an opportunity to gain insight into the metabolic behavior of individual cell types. Here, we estimated the availability of nutrients and oxygen within the tumor microenvironment using concentration measurements from blood together with a metabolite diffusion model. In addition, we developed an approach to efficiently apply enzyme usage constraints in a comprehensive metabolic model of human cells. The combined modeling reproduced severe hypoxic conditions and the Warburg effect, and we found that limitations in enzymatic capacity contribute to cancer cells' preferential use of glutamine as a substrate to the citric acid cycle. Furthermore, we investigated the common hypothesis that some stromal cells are exploited by cancer cells to produce metabolites useful for the cancer cells. We identified over 200 potential metabolites that could support collaboration between cancer cells and cancer-associated fibroblasts, but when limiting to metabolites previously identified to participate in such collaboration, no growth advantage was observed. Our work highlights the importance of enzymatic capacity limitations for cell behaviors and exemplifies the utility of enzyme-constrained models for accurate prediction of metabolism in cells and tumor microenvironments.

INTRODUCTION

The tumor microenvironment (TME) consists of many different cell types that share a common influx of metabolites from the blood, and its leaky blood vessels make diffusion the main mechanism for transport of metabolites and oxygen from the blood to the cells.^{1–4} Furthermore, the blood vessel density is unevenly distributed, leading to a highly variable availability of nutrients and oxygen, where hypoxic and even necrotic regions tend to be more common in the center of the tumor.⁵ Cancer cells residing in the TME are generally thought to strive for proliferation, and the metabolic behavior that is optimal for maximizing growth will vary with the availability of metabolites. Such known behaviors are lactate export (the Warburg effect⁶), which is shared with other highly proliferative cells such as expanding T cells,⁷ and “glutamine addiction”,⁸ where the tricarboxylic acid (TCA) cycle is preferentially fed with glutamine. Furthermore, bypass of mitochondrial complex I can be used to increase ATP production at a lower substrate efficiency in cells with high ATP demand,⁹ although such a behavior has not explicitly been proven to exist in cancers.

Collaboration scenarios between different “healthy” cell types and tumor cells, where cells such as fibroblasts and macrophages are thought to be exploited by the cancer cells, has been a topic of great interest for the past decade.^{10–13} Specifically, these cells are thought to regulate the tumor microenvironment through signaling as well as provide nutrients such as lactate and pyruvate to the tumor cells to enhance their growth. Partial evidence of the latter has been presented, where cancer-associated fibroblasts have been observed to secrete metabolites that were taken up by cancer cells,¹² but there is no solid evidence that such collaboration is directly beneficial for growth.

The study of collaborative metabolism in tumors is difficult for two reasons: (1) It is challenging to create a realistic *in vitro* representation of the tumor microen-

vironment using cell lines. For example, the influx of oxygen and all other metabolites from blood needs to be controlled and constant over time. (2) It is difficult to measure metabolite uptake rates *in vivo* for all metabolites, although isotope labeling has successfully been employed for single metabolites.¹⁴ An alternative approach is therefore needed.

Genome-scale modeling¹⁵ of human metabolism using flux balance analysis (FBA) involves performing *in silico* analyses of a reaction network under steady-state conditions and has been used to investigate metabolism in, for example, muscles,⁹ tumors,¹⁶ and Alzheimer's disease.¹⁷ A recent approach is the GECKO framework,¹⁸ which enables the integration of enzyme kinetic data with genome-scale metabolic models to simulate more physiologically meaningful flux distributions even when metabolite exchange rate data are limited.

In this project, we developed a diffusion model for constraining metabolite uptake rates in tumors and applied enzyme usage constraints to a human genome-scale model to simulate tumor metabolism. Our metabolic models showed that glucose and oxygen are the most limiting substances for growth. Furthermore, our models predicted glutamine addiction, where fueling the TCA cycle with glutamine can yield more ATP since it requires less enzyme usage per ATP produced. In addition, we observed that metabolic collaboration between cancer-associated fibroblasts (CAFs) and cancer cells likely has a very small effect on tumor growth.

RESULTS

Simulation of tumor cell growth

We used the genome-scale metabolic model Human1¹⁹ enhanced with enzyme constraints to model the metabolism of cancer cells in the tumor microenvironment. The model was manually curated (materials and methods, Table S1) and a non-growth-associated maintenance (NGAM) requirement was added as an ATP cost of 1.833 mmol gDW⁻¹ h⁻¹ derived from literature.^{20,21} We developed a lightweight version of the GECKO toolbox,^{18,22} called GECKO Light (Note S1), which, similar to GECKO, constrains the total metabolic enzyme usage based on k_{cat} values from the BRENDA database.²³ The main improvement in GECKO Light is a substantially decreased execution time (~2 min on a standard laptop computer) and a smaller generated model size.

We assumed that diffusion is the dominant mechanism for influx of metabolites into tumors²⁴ and developed a diffusion model that uses the metabolite concentrations in blood and their diffusion coefficients to estimate the metabolite uptake constraints (Figure 1A, Note S2). Instead of estimating absolute uptake constraints, which are expected to vary across cells depending on many factors such as distance to capillaries, we estimated relative uptake constraints according to

$$U_i = aD_i c_{b,i}$$

where U_i is the estimated upper bound for the uptake flux of metabolite i , a is a proportionality constant that is inversely related to the distance to the closest capillaries, D_i is the diffusion coefficient for metabolite i , and $c_{b,i}$ is the concentration of metabolite i in the blood. While the value of a varies across cells and is difficult to estimate for a given cell, its value will be the same for all metabolites within the same cell, which makes it possible to investigate the metabolism at different pseudodistances from blood

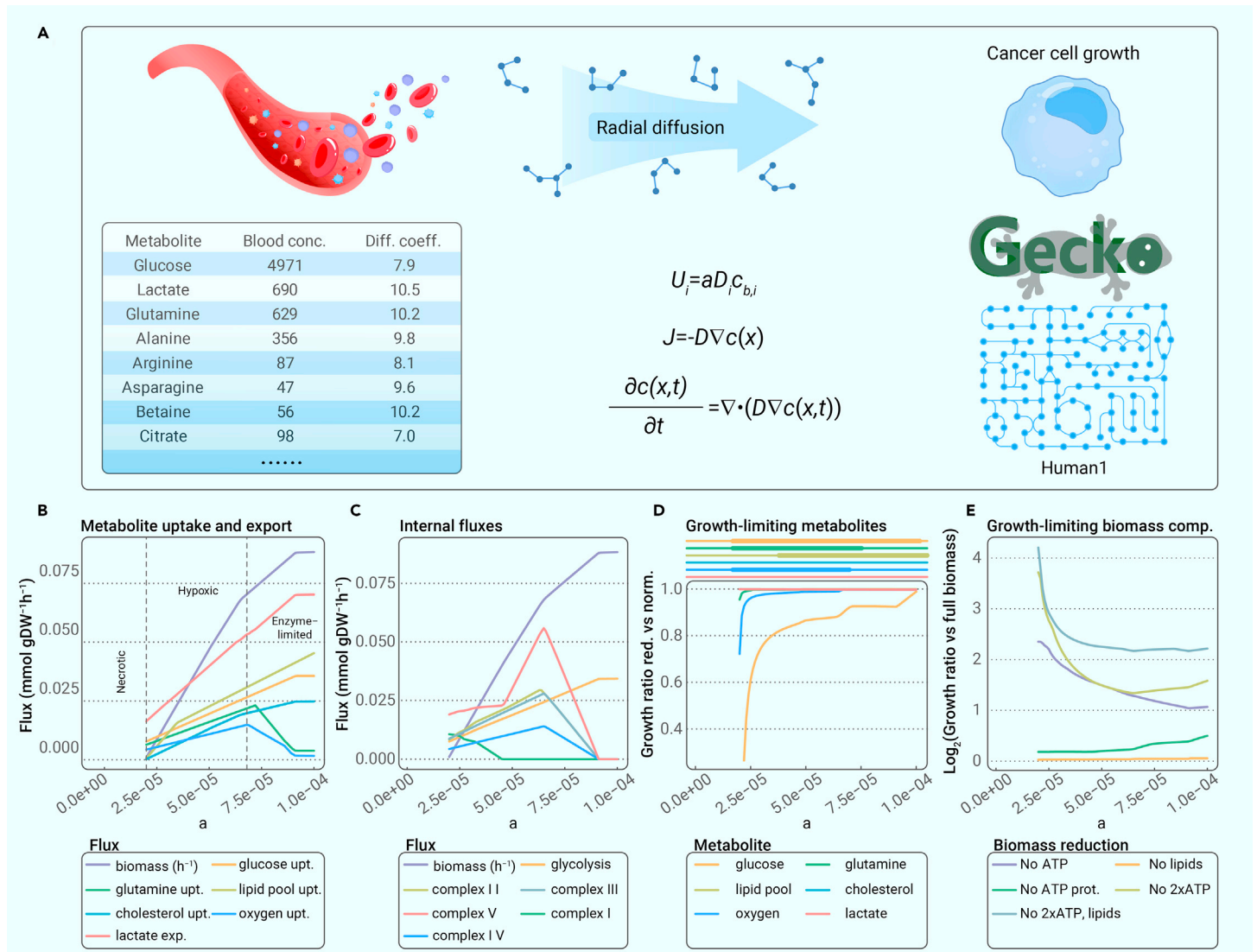


Figure 1. Modeling cancer cell growth in the tumor microenvironment (A) Modeling setup. The metabolite uptake bounds are estimated from a diffusion model based on metabolite concentrations in blood and their diffusion coefficients. In addition, enzyme usage constraints are added to the model using GECKO Light. (B) Simulated specific growth rate (biomass production) and metabolite exchange rates for different values using the diffusion model. Some fluxes are scaled to enable visualization of all metabolites in the same figure: glucose upt.: 0.01, glutamine upt.: 0.05, Lipid pool upt.: 0.2, cholesterol upt.: 10, oxygen upt.: 0.1, and lactate exp.: 0.01. (C) Internal fluxes in the same simulations as (B). The glycolytic flux is multiplied by 0.01 and the fluxes through the complexes are multiplied by 0.1. (D) Growth dependence of metabolites. (Top) A thick line indicates a required uptake rate of at least 95% of a metabolite for maintaining growth, as determined by flux variability analysis (FVA). (Bottom) The simulated specific growth rate of the model is compared with that when the maximum uptake rate of a single metabolite is reduced to 90%. (E) Change in specific growth rate when removing parts of the biomass reaction. In all cases, the model is optimized for growth. "No ATP prot." refers to removal of the ATP cost from polymerizing amino acids into proteins, while "No 2xATP" refers to having both the protein generation ATP cost and the growth-associated maintenance (GAM) ATP cost removed from the biomass reaction. For "No 2xATP, lipids," the consumption of lipids has also been removed in addition to the other two factors. See Figure S5 for details.

vessels. In addition, we developed a blood flow model, where the blood flow into the tumor rather than diffusion was assumed to be the limiting factor for delivery of metabolites to cells and the influx was modeled to be directly proportional to the contents of the blood (Note S3).

We retrieved blood plasma measurements from several sources to form a collection of 69 metabolites with associated concentration values^{25–28} (Note S4; Tables S2, S3, and S4; Figure S1). Lipids were grouped into sterols (represented by cholesterol) and other lipids (fatty acyls, glycerolipids, glycerophospholipids, and sphingolipids, represented as a mix of fatty acids). For oxygen we used the concentration of free oxygen, which can diffuse, excluding roughly 98% of the total concentration in blood that is bound to hemoglobin.

We collected 18 metabolite diffusion coefficients from literature,^{29–31} and estimated the remaining coefficients using a linear model based on molecular mass³² (Figure S2, Note S5; Table S4). Lipids are not soluble in water and are therefore transported together with albumin or a lipoprotein, or potentially as droplets. These are large particles that do not diffuse well, and we modeled the diffusion of these particles by using the diffusion coefficient of albumin for all lipids.

We first simulated the tumor cell growth for different values of the proportionality constant a using FBA (Figure 1B). For small values of a (greater distances from capillaries), the model cannot produce enough ATP to uphold cell maintenance requirements, resulting in necrosis. For higher values of a , we see a hypoxic region where the cells can survive and grow but are limited in growth by availability of nutrients and oxygen, followed by a region where the enzyme constraints become the major limiting factor for growth. At small values of a , the limited oxygen supply was primarily used for oxidative phosphorylation, which provided much ATP per substrate. However, when more nutrients were available a larger specific growth rate could be reached by allocating more of the enzymatic capacity to glycolysis instead of to these large and slow enzyme complexes. The simulation predicts an early bypass of complex I, while the remaining electron transport chain (ETC) complexes remain active at a wider a range (Figure 1C), which is consistent with a previous study where complex I bypass was described to increase ATP yield at the cost of a lower substrate efficiency.⁹ The model exhibits an extreme behavior where oxidative phosphorylation is completely shut down for large values of a . In practice, such a behavior is generally not observed. Eighty percent of the ATP has for example been reported to be

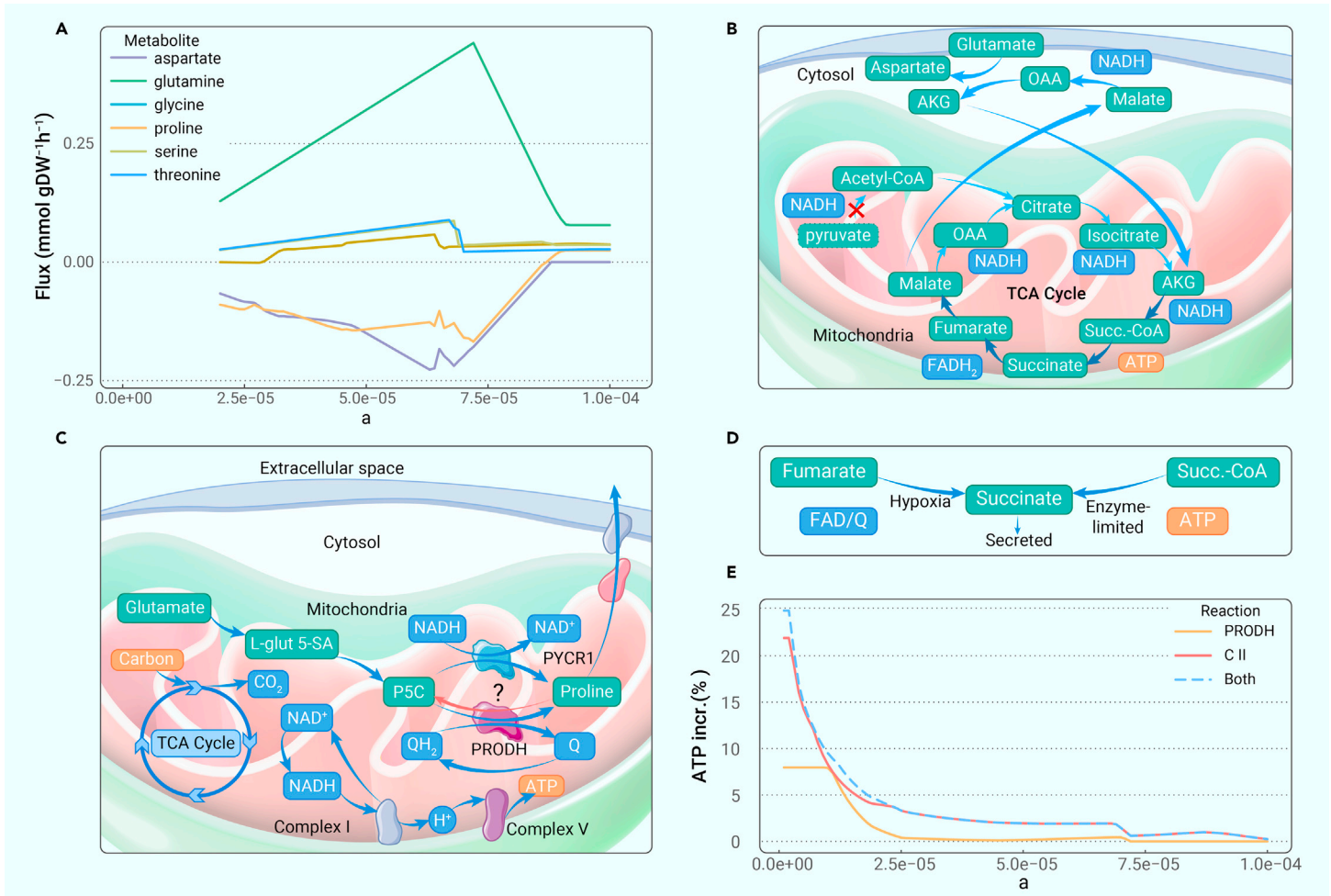


Figure 2. Predicted amino acid metabolism in simulation of cancer cell growth (A) amino acid fluxes from the simulation used in Figure 1B. Only amino acids with high predicted fluxes are shown. Positive fluxes correspond to cellular uptake and negative to export. (B) Predicted “glutamine addition” mechanism. When OXPHOS is limited by its high enzyme usage, the TCA cycle flux is limited because there is no way for the cell to oxidize the produced NADH/FADH₂. Any processes that can either limit the production of NADH/FADH₂ while running the TCA cycle or increase oxidation of NADH/FADH₂ will thus lead to a possibility to increase the flux through the TCA cycle, and thereby increase the ATP production. Feeding the TCA cycle from glutamate instead of pyruvate shortcuts the TCA cycle, reducing two less NAD⁺ molecules to NADH, while also preserving pyruvate that can be converted into lactate to oxidize another NADH molecule. Darker arrows indicate larger fluxes. (C) Proline metabolism. Generation of proline can help to increase flux through the TCA cycle as well as bypass complex I and thereby increase ATP production. The model predicts two mechanisms for this purpose: (1) Oxidation of NADH through any of the PCYR enzymes (complex I bypass); and (2) by running PRODH in reverse (non-favorable thermodynamically, as indicated by the question mark). The latter would increase flux through complex I, since Δ¹-pyrroline-5-carboxylate (P5C) would replace oxygen as the electron acceptor in the electron transport chain, and could therefore in theory increase ATP production by consuming excess NADH and increasing flux through complex V. The brown arrow represents the proline cycle, where PRODH is run in the thermodynamically favorable direction together with the use of a PCYR enzyme, which forms a cycle that can potentially move electrons from NADH to ubiquinone, thereby bypassing complex I without exporting proline. (D) Succinate export. Allowing succinate export yields two advantages: (1) Complex II can be run in reverse, which highly resembles the effect from running PRODH in reverse; and (2) the TCA cycle can be stopped at succinate, reducing production of NADH and FADH₂, which enables the cell to run the TCA cycle more rounds and thereby produce more ATP. (E) Increase in ATP production from allowing the mechanisms explained in (C) (PRODH in reverse) and (D) (complex II in reverse). The model is optimized for ATP production, the NGAM is set to 0, and the metabolite uptake bounds are determined by the diffusion model.

generated by oxidative phosphorylation in some highly proliferative cells,³³ but regardless of this limitation the model can still serve as a means to understand observed metabolic behaviors.

Next, we evaluated which metabolites were limiting for growth (Figures 1D and S3). Glucose and oxygen were by far the two most important metabolites for growth, followed by a smaller effect from glutamine, while the other limiting metabolites had a small effect (Figure 1D). Neither lactate nor albumin was limiting for growth since neither of them can give a positive ATP contribution without using oxygen. Notably, some metabolites such as lipids were limiting for growth, but the effect is marginal; other substrates can be used in their place with an almost negligible loss in growth rate. For the hypoxic range, free access to oxygen yielded a clear increase in growth (Figure S4). We repeated our simulations while removing different components of the biomass reaction to investigate which cellular processes are limiting for growth, which showed that ATP production was the main limiting factor (Figures 1E and S5). To assess the sensitivity of our results, we repeated our analyses with the growth-associated ATP cost and NGAM reduced to 50% and 25% of their original values, which confirmed that glucose and oxygen were still the most important contributors to growth (Figure S6). Since hypoxia is less common in the healthy body compared with

tumors, we also identified five reactions that were important for growth in hypoxia but not in normoxia as potential targets (Table S5). However, all such reactions were related to oxidative phosphorylation and are likely not suitable targets. The lack of such targets is likely caused in part by redundancies in the full metabolic network, and to identify more targets in future studies, cell-type-specific reaction networks derived from for example single-cell RNA-seq data from specific cancer types could be used.

The blood flow model produced similar results, although with a substantially smaller hypoxic region (Figure S7). The major difference is a higher availability of lipids and oxygen, which increases oxygen usage at small *a* values.

Amino acid metabolism

The diffusion model predicted large uptake fluxes of glutamine, glycine, serine, and threonine, and export of proline and aspartate, while the rest of the amino acids were consumed at lower fluxes, primarily for protein synthesis (Figures 2A and S8). The model did not predict glutamate secretion, which is observed in some cancer cell lines.^{34,35} While glutamate export has previously been linked to nucleotide synthesis,³⁵ our model does not predict any advantage of such a behavior in the TME.

Table 1. ATP production from amino acids compared with lactate in different settings

Substrate	No O ₂	Low O ₂	Low O ₂ PRODH	Low O ₂ Sc. Exp.	Enz. Lim.	Enz. lim. Sc. Exp.
Lactate	0	5	5	6.5	0.0938	0.0938
Aspartate	0	5	6.5	12.9	<u>0.0937</u>	<u>0.0937</u>
Glutamine	0	5	10	9.1	0.1042	0.1281
Glycine	0	5.7	5.7	8	<u>0.0929</u>	<u>0.0929</u>
Proline	0	<u>4.7</u>	<u>4.7</u>	<u>4.9</u>	0.0944	0.0952
Serine	3.33	9	9	13.0	0.0945	0.0945
Threonine	3.33	9	9	12.5	<u>0.0931</u>	<u>0.0931</u>
Alanine	0	5	5	6.5	<u>0.0932</u>	<u>0.0932</u>
Arginine	0	5	12.3	<u>5.4</u>	0.0943	0.0949
Asparagine	0	5	6.5	12.9	0.0940	0.0944
Cysteine	0	5	5	<u>5.9</u>	<u>0.0927</u>	<u>0.0927</u>
Glutamate	0	5	10	9.1	0.1020	0.1213
Histidine	0	5	11.1	7.3	<u>0.0927</u>	<u>0.0927</u>
Isoleucine	0	5	5	5	<u>0.0922</u>	<u>0.0922</u>
Leucine	0	5	5	5	<u>0.0917</u>	<u>0.0917</u>
Lysine	0	<u>4.8</u>	<u>4.8</u>	<u>4.8</u>	<u>0.0918</u>	<u>0.0918</u>
Methionine	0	5	5	5	<u>0.0830</u>	<u>0.0830</u>
Phenylalanine	0	5	5	5	<u>0.0900</u>	<u>0.0900</u>
Tryptophan	0	<u>3.8</u>	<u>3.8</u>	<u>3.8</u>	<u>0.0877</u>	<u>0.0877</u>
Tyrosine	0	5	5	5	<u>0.0916</u>	<u>0.0916</u>
Valine	0	5	5	5	<u>0.0917</u>	<u>0.0917</u>

The table shows the maximum ATP production ($\text{mmol} \cdot \text{gDW}^{-1} \cdot \text{h}^{-1}$) given a maximal uptake of $10 \text{ mmol} \cdot \text{gDW}^{-1} \cdot \text{h}^{-1}$ of a single substrate and varying oxygen availability. Columns: "No O₂": No oxygen uptake is allowed, negligible effects from enzyme constraints. "Low O₂": Oxygen uptake constrained to $1 \text{ mmol} \cdot \text{gDW}^{-1} \cdot \text{h}^{-1}$ (which is not enough to fully oxidize any of the substrates), negligible effects from enzyme constraints. PRODH in reverse and succinate export are both blocked. "Low O₂ PRODH": Same as "Low O₂", but with PRODH in reverse enabled. "Low O₂ Sc. Exp.": Same as "Low O₂", but with succinate export enabled. "Enz. Lim.": The total available enzyme pool is constrained to a low value (0.001 g/gDW). Succinate export is blocked. In practice, this also means that the reverse PRODH reaction will not be used. "Enz. Lim. Sc. Exp.": Same as "Enz. Lim." but allowing for succinate export. Bold and italic text correspond to a higher ATP production compared with lactate and underlined text to a lower flux. While comparison to using pyruvate as fuel may at first seem more relevant since pyruvate is the alternative fuel for the TCA cycle coming from glycolysis, the pyruvate will be exported as lactate if not used, making comparison to lactate fairer from a redox perspective.

The genome-scale metabolic model can be used to study in detail the most optimal amino acid metabolism in different conditions, where the NADH/FADH₂ oxidation ratio in the ETC plays an important role. For fast-growing cells limited by enzymatic capacity, the ratio should be as low as possible to maximize ATP production (complex I bypass), while the ratio should be as high as possible in oxygen-deprived conditions (Note S6). Cells can utilize amino acids in different ways to manipulate this ratio.

"Glutamine addiction" is a well-known cancer trait,^{8,36–38} where glutamine, the most highly abundant amino acid in blood, is used instead of glucose-derived pyruvate to feed the TCA cycle. Indeed, the model predicts such behavior. Glutamine is converted to glutamate and enters the TCA cycle, producing three fewer NADHs per cycle than if pyruvate is used, resulting in increased ATP production, aspartate production, and reduced reaction oxygen species (ROS) generation (Figure 2B and Note S6). While aspartate export was not observed in a cell culture experiment of the NCI-60 cell lines,³⁹ aspartate production from glutamine has

been shown to exist in pancreatic cancer cell lines using labeling experiments,⁴⁰ and is likely disposed of using alternative pathways (Note S6). Glutamine is also known to be important for immune cells such as T and NK cells, possibly for the same purpose, i.e., to increase ATP production. However, due to the predatory uptake of glutamine by cancer cells, glutamine is often depleted in the TME, making this behavior a weakness in the TME.⁴¹

Proline secretion has been observed in cell line experiments^{35,39} and the model predicts two different mechanisms by which proline production from glutamate can help to increase ATP production: NADH oxidation through PYCR1 (or related enzymes), which can also form the "proline cycle"⁴² with PRODH to bypass complex I, or by running PRODH in reverse, enabling the use of complex I without oxygen (Figures 2C and S9). The latter has to our knowledge not been reported, and it is unclear if such a reaction is thermodynamically favorable *in vivo*, although it has been shown possible in enzyme assays outside cells.⁴³ In addition, allowing export of succinate enables (1) the reversal of complex II, which catalyzes a similar reaction and has been shown to run in reverse in mammals (Figure 2D)^{44,45}; and (2) interruption of the TCA cycle, thereby bypassing complex II while running succinyl-CoA synthetase to produce ATP (Figure 2D). The model predicts these two effects to be even more beneficial than PRODH in reverse under hypoxia, although all effects combined yield the highest benefit (Figures 2E and S10).

To investigate the behavior of each amino acid in more detail, we simulated access to a fixed influx of a single metabolite and optimized for maximum ATP production, comparing the use of each amino acid as the sole carbon source to that of using lactate under different conditions (Table 1). A few amino acids could be used to generate ATP without oxygen, such as serine, which as part of the serine, one-carbon cycle, glycine synthesis (SOG) pathway has been shown to support ATP production and cancer cell proliferation.⁴⁶ However, like lactate most amino acids were not useful for producing ATP in that condition. In hypoxia, without enzyme constraints, PRODH reversal, and succinate export, some benefits to ATP production could be observed from using amino acids as substrates compared to lactate. Allowing reversal of PRODH increased the ATP yield from several amino acids dramatically compared with lactate, and a similar effect was observed from allowing export of succinate, although the benefits varied substantially across different amino acids. Under enzyme-limited conditions, both with and without succinate export, glutamine and glutamate were the carbon sources giving the highest ATP yield, consistent with the strategy depicted in Figure 2B and the glutamine addition results reported previously.⁸ Under these conditions, complex I was bypassed, which resulted in a several times higher TCA cycle flux for glutamine and a higher total ATP production despite having lower complex V activity (Figure S11). Comparing these results to existing literature, a previous study showed that glutamine is used to feed the TCA cycle both in hypoxic and normoxic conditions,³³ which is consistent with our results assuming either use of PRODH in reverse or succinate export in hypoxia.

Simulation of cell-type collaborations in the TME

A topic of recent interest is whether non-cancerous cells in the TME assist cancer cells metabolically by providing them with resources that are advantageous for growth. It has for example been proposed that CAFs can provide cancer cells with metabolites such as lactate and pyruvate,^{10,47} and it could also be beneficial if tumor-associated macrophages (TAMs) could consume dead cells and cellular debris and produce nutrients for the cancer cells. We sought to investigate these collaboration scenarios in more detail using our diffusion modeling approach. We again constrained metabolite uptake from blood but built a more complex metabolic model consisting of three cell types: cancer cells, fibroblasts, and other cells, where the latter represent cells that are not expected to provide resources to the cancer cells, for example immune cells (Figure 3A). The exchange of metabolites between the cell types was controlled by providing separate compartments for the interstitium around the fibroblasts and other cells (Figure 3B).

The extracellular matrix (ECM) consists mainly of collagens and glycosaminoglycans (GAGs) and is produced by the fibroblasts. Both the composition of the ECM and the fraction of the total tumor dry weight that the ECM constitutes vary largely across tumors. We assumed an ECM composition of 80% collagen (represented by collagen I) and 20% GAGs (represented by heparan sulfate) to reduce the number of flexible parameters. We varied the total ECM fraction of the tumor dry weight and the fraction of each cell type, since these parameters vary substantially between individual tumors. The different modeling configurations are listed in Table 2.

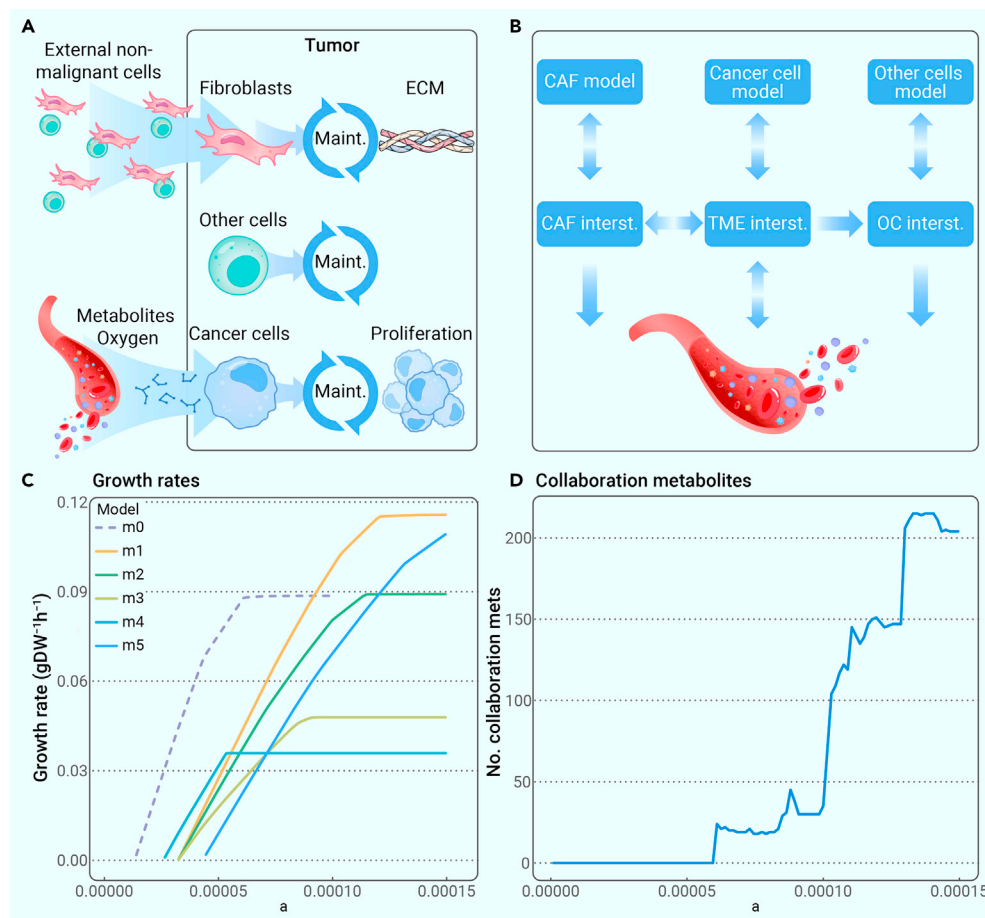


Figure 3. Collaboration between cell types in the TME (A) Modeling setup of the tumor microenvironment. The model contains three cell types: Fibroblasts (CAFs), cancer cells, and other cells (e.g., different types of immune cells). Non-malignant cells are imported to the tumor, resulting in a total biomass reaction consisting of cancer cell growth and extracellular matrix construction. All cell types have the same metabolic network (the curated full Human1 model) with the same NGAM value as used previously, but the fibroblasts were extended with reactions for building the extracellular matrix. (B) Compartment setup to enable limitation of cell-type collaboration to certain scenarios. The fibroblasts (CAF) and other cells (OC) have their own model compartments for the tumor interstitium. While metabolites can (depending on the simulation) be transported from the CAF interstitium to the cancer cell interstitium, such transport is not possible from the OC interstitium. (C) Effect of collaboration on the specific growth rate of the tumor. (D) Number of collaboration metabolites between fibroblasts and cancer cells as a function of *a*, measured for the *m2* model. We identified in total 233 potential collaboration metabolites across the range of explored *a* values.

It directly follows from the first law of thermodynamics that CAFs cannot increase the total energy available in the TME unless they actively import or utilize material from outside the TME, which is a case not investigated here but briefly addressed later in this section. In cases where the growth is limited by nutrient availability only, there is therefore nothing the CAFs can do to help the cancer cells metabolically, given that the cancer cells behave optimally and thereby express the enzymes needed to utilize the available nutrients. However, as nutrient availability increases and the growth becomes limited by enzymatic capacity, it may be possible for the CAFs to increase growth by providing additional enzymatic capacity to produce metabolites that are more optimal for cancer cell growth.

We first simulated the specific growth rates of the models with different parameter values for ECM and cell-type fractions (*m1*–*m5*) and compared them with that of the *m0* model (Figure 3C). A high fraction of fibroblasts penalized growth at low values of *a* due to the additional ATP costs from NGAM, while the extra enzymatic capacity supplied by a high fibroblast fraction imparts a growth advantage for high *a* values. We also observed that the cost of building the ECM was not negligible in our simulations. However, the addition of ECM to the model objective function did not substantially alter which metabolites were limiting for growth relative to the *m0* model, despite a high protein content (Figure S12A).

Next, we developed an algorithm to identify potential *collaboration metabolites*, defined as metabolites being exported from fibroblasts and imported into cancer cells, for all values of *a* (Figure 3D; Table S6). As expected, the number increased with increasing *a*, where the enzymatic capacity becomes increasingly limiting for growth. However, many of these collaboration metabolites, such as ATP, are not physiologically meaningful and have not been reported as being exported from fibroblasts. When limiting the collaboration metabolites to those reported in literature (lactate, pyruvate, free fatty acids, glutamine, ketone bodies, and alanine^{47,48}), and comparing with the model in which collaboration is blocked, the growth difference was negligible (Figure S12B). Although co-injection of tumor cells with fibroblasts in mice can lead to an increased specific growth rate,⁴⁹ we reason that this effect is likely caused by signaling effects or potentially

the given set of collaboration metabolites were therefore evaluated in our simulations. Other cell types, for example myeloid-derived suppressor cells (MDSCs), have a different objective and are additionally known to export ROS and reactive nitrogen species (RNS).⁵² We therefore extended the set of collaboration metabolites with H₂O₂, which was the only ROS/RNS species identified as a collaboration metabolite, and used the *m1* model with a negligible ECM fraction and allowed the CAFs to serve as MDSCs, but only negligible changes to the growth rate were observed (Figure S12C).

Although the “other cells” cannot provide the cancer cells with resources in the simulation, they can still minimize their negative impact on growth, which we have termed passive collaboration. We simulated this behavior by maximizing tumor growth using the *m6* model, resulting in ATP generation based on enzyme-demanding processing such as oxidative phosphorylation in the other cells (Figure S12D). The question is whether the other cells, representing for example immune cells, are helpful for cancer growth, or if the relationship is more competitive. TAMs have, however, been reported to switch to oxidative phosphorylation when exposed to lactate, which could be seen as a sign of such passive collaboration.⁵³

TAMs are reported to have many metabolic roles in the TME, including both passive and active collaboration,^{53,54} which would give them the same role as other cells or fibroblasts in our simulation. Macrophages, however, also have an additional function where they can scavenge dead cells and convert them into metabolites useful to other cells. To investigate to what extent this function can support tumor growth, we used the *m0* model and assumed that 10% of the tumor cells produced from growth die. We assumed that the maximum amount of additional nutrients provided is thus the metabolites that constitute 10% of the produced biomass (materials and methods), excluding the ATP cost consumed during growth (Table S7). There is a clear growth advantage, which is larger for small *a* values, but the overall effect is relatively small even though no maintenance or metabolite conversion cost is included for the macrophages (Figure S12E). The reason for the limited utility of such substrates is mainly that in nutrient-deprived conditions, all oxygen is already being consumed, and oxygen is required to generate ATP from most scavenged resources. A similar theory to

Table 2. Different model configurations used in the simulations

Model	Cancer cell fraction	Fibroblast cell fraction	Other cells fraction	ECM fraction
<i>m0</i>	1	0	0	0
<i>m1</i>	0.6	0.2	0.2	0.01
<i>m2</i>	0.6	0.2	0.2	0.25
<i>m3</i>	0.6	0.2	0.2	0.5
<i>m4</i>	0.75	0.05	0.2	0.25
<i>m5</i>	0.45	0.35	0.2	0.25
<i>m6</i>	0.9699	0.0001	0.03	0.00001

The *m0* model contains only tumor cells and is identical to the model used in Figures 1 and 2. ECM fraction represents the weight fraction of the total objective being maximized that the ECM constitutes.

Permissions: Not applicable.

that of scavenging dead cells by macrophages concerns catabolism of organelles and apoptosis in CAFs to provide additional nutrients,¹⁰ where CAFs could then for example be recruited to the tumor. Such a case represents an influx of extra material from outside the TME and could provide a small growth advantage similar to the macrophage case but also suffers from the same problem of oxygen limitation and is likely of limited importance.

We also performed these simulations with the blood flow model. In contrast to the diffusion model, a small (negligible) collaboration benefit was observed from collaboration using literature metabolites (Figure S14A), and the macrophage collaboration utilizing dead cells was more beneficial in this case (Figure S14B). However, the benefits were still small, confirming our previous conclusion that these metabolic collaboration scenarios are not important for optimizing cancer growth.

DISCUSSION

During the past decade, metabolic collaboration between cell types in the tumor microenvironment has been thought to increase tumor growth,^{10–13} although experimental evidence is lacking due to difficulties in measuring and controlling fluxes into and between cell types in an experimental setup. As an alternative, we used genome-scale modeling of the human intracellular metabolic network and integrated measurements of metabolite concentrations in blood and metabolite diffusion coefficients to estimate the maximal increase in specific growth rate from such collaborations at different levels of hypoxia.

The major insight gained from this study was that the metabolic collaboration scenarios between cell types proposed in the literature^{10–13} have a negligible effect on tumor growth. Although fibroblasts have been shown to export lactate⁵⁵ and some tumors have been shown to take up lactate,⁵⁶ no direct evidence is presented that such a behavior is beneficial for growth. In addition, scavenging of dead cells by macrophages or other forms of cell catabolism such as catabolic behavior of fibroblasts¹³ only yield a marginal benefit to growth due to a shortage of oxygen. However, the signaling interplay between stromal cells and cancer cells is most likely important for changing the metabolism of tumor cells.¹³

An additional insight gained from this study is that ATP production is most likely the limiting factor for growth of cancer cells in the TME, and that the ATP production is limited by either lack of nutrients and oxygen, by limitations in enzymatic capacity, or a combination of both. Other factors not considered in our model may also play a role in limiting growth, such as extensive ROS production or low pH in the interstitial fluid. The high demand for ATP is implicitly supported by literature; most of the carbon from glucose and glutamine imported by cells is not incorporated into the biomass,⁵⁷ suggesting that it is used for ATP production.

The third important insight from this study concerns the use of amino acids for energy production and the varying ways in which the TCA cycle can be used. Our model, for example, predicts glutamine addiction^{8,58,59} and gives plausible explanations to why glutamine is a more optimal substrate than pyruvate for the TCA cycle at different levels of hypoxia. We also predict potential benefits from running PRODH in reverse in hypoxia, although it is unclear if such a reaction

is thermodynamically favorable, and predict a potential 20% increase in ATP production from running mitochondrial complex II in reverse, which has been shown to take place both in cell lines and *in vivo*.⁴⁵

Although we have shown that direct metabolic collaboration between cells in the TME is not useful for increasing the growth rate of tumor cells, metabolic interactions in the TME are still important. A recent example is the ability of the oncometabolite D-2-hydroxyglutarate (D-2HG) to block lactate dehydrogenase.⁶⁰ The acquisition of mutations in isocitrate dehydrogenase by cancer cells to produce this oncometabolite is at first counterintuitive—we predict that blocking lactate production reduces the growth rate of cancer cells. However, as shown in the aforementioned study, it also blocks lactate production in T cells, which impairs cytotoxicity and thereby increases the survival of cancer cells. Likewise, as we have discussed in this study, depletion of glutamine can potentially reduce the toxicity of T cells.⁴¹ Metabolic drivers in cancer therefore need not necessarily increase growth but can also provide other benefits.

Altogether, our results suggest that there is no benefit for tumors to develop a metabolic collaboration between cell types. Such metabolic collaborations may still exist, where a cell type can complement deficiencies in tumor cells, which has for example been observed during glutamine deprivation.⁶¹ However, such a configuration does not improve growth beyond what activating the needed pathway directly in the tumor cells would yield. Our modeling approach allows for investigation of the optimal behavior for cells in the TME under the assumptions that the diffusion and blood flow models are based upon. While these models are approximations of the true influx of metabolites into tumors, they can still serve well for future investigations of different metabolic phenomena in the tumor microenvironment, and with a systems biology approach unravel questions that are difficult to address using experiments.

MATERIALS AND METHODS

Details about the materials and methods used are available in Note S7.

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ACKNOWLEDGMENTS

We thank Stig Larsson for reviewing the diffusion model and Gunjan Purohit for help with investigating amino acid metabolism.

This work was supported by funding from the Knut and Alice Wallenberg Foundation, Grant 2015-0279 (J.N.) and in part by NIGMS, National Institutes of Health, Grant R01GM13264 (D.B.).

AUTHOR CONTRIBUTIONS

J.G., J.R., and J.N. planned the project. J.G. and A.H. developed the diffusion model. J.G., F.R., D.B., S.P., and O.D. investigated and drew conclusions about amino acid metabolism. J.G. wrote all the software and made all analyses and figures. J.G. wrote the draft manuscript. All authors reviewed and edited the manuscript. J.R. and J.N. supervised the project. J.N. acquired funding for the project.

DECLARATION OF INTERESTS

The authors declare no competing interests.

SUPPLEMENTAL INFORMATION

It can be found online at <https://doi.org/10.1016/j.xinn.2024.100583>.