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Tumor-on-chip platforms for transport phenotyping: decoding CAF-driven barriers to drug delivery

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Physical barriers within solid tumors constitute a fundamental but often overlooked mechanism of therapeutic resistance, contributing to the poor success rate of cancer drug translation. Therapeutic molecules often fail to reach their intended targets due to mass-transport limitations imposed by the remodeled, spatially heterogeneous tumor microenvironment (TME). Cancer-associated fibroblasts (CAFs) drive dynamic remodeling of the extracellular matrix (ECM), generating local variations in stiffness, porosity, and intrinsic permeability that, together, shape evolving transport phenotypes that govern drug accessibility. We focus on mechanical pathways of stromal mechanotransduction, tracing the sequence from CAF activation through ECM remodeling, to barrier formation, and show how these processes collectively govern therapeutic outcomes. We also evaluate advanced microfluidic and tumor-on-chip (ToC) platforms that reproduce stromal heterogeneity under controlled conditions, mimicking tissue architecture, transport behavior, and therapeutic response. By enabling patient-specific profiling of CAF-driven transport phenotypes, these systems demonstrate that transport barriers are not fixed obstacles but dynamically modifiable therapeutic targets. “Transport phenotyping” could complement genomic profiling in clinical oncology by integrating heterogeneity, biophysics, and precision medicine, potentially transforming personalized treatment strategies for patients whose tumors remain refractory to current therapies.

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1. Introduction

Contemporary cancer therapeutics face a fundamental paradox: despite unprecedented molecular characterization of oncogenic processes and the sophisticated development of targeted agents, recurrent and metastatic solid tumors remain largely refractory to treatment. This paradox contributes to poor patient survival outcomes. Annual approval statistics show that new chemical entities often display remarkable preclinical efficacy, yet around 95% fail to achieve meaningful survival benefits in clinical trials, with most failures occurring in phase II and III due to lack of effectiveness.^{1,2} This pattern of early discovery success but late-stage attrition cannot be explained solely by biological complexity or tumor heterogeneity and plasticity. An increasingly recognized obstacle lies in the physical architecture of solid tumors: a dense, fibroblastic (desmoplastic) stroma that prevents drugs, from small molecules to antibodies and nanoparticles, from reaching malignant cells.³ Intrinsic drug resistance reflects cell-autonomous mechanisms such as mutations in molecular targets, activation of efflux pumps, or

enhanced DNA repair pathways. Extrinsic resistance arises from the tumor microenvironment (TME), which acts as both a physical and biochemical barrier to treatment. While genetic and molecular resistance mechanisms have been extensively reviewed,^{4,5} we focus here on the physical microenvironment, notably the extracellular matrix (ECM), as a critical determinant of extrinsic, transport-mediated resistance.‡ This stromal barrier is not generated by tumor cells themselves but by the cellular components of the TME, cancer-associated fibroblasts (CAFs), through ECM remodeling and mechanical stress. Tumor heterogeneity and cell plasticity occur across all solid cancers and vary not only between patients but also among regions of the same lesion.⁶ Beyond cellular variability, the ECM undergoes continuous remodeling, generating spatially and temporally heterogeneous stiffness, porosity, and intrinsic permeability that govern drug accessibility.^{7,8} Here, we use the term intrinsic permeability (k) to denote the matrix property of the ECM, distinguishing it from hydraulic conductivity, which additionally depends on fluid viscosity and is often used inconsistently in the biological literature. Such remodeling is dynamic, nonlinear, and shaped by CAF heterogeneity and biochemical-mechanical feedbacks.^{9–11} CAF-driven matrix remodeling thus shapes evolving transport landscapes that determine whether therapeutic agents and immune cells can reach malignant cells.^{12,13}

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‡ Hereafter, the term “transport” refers to molecular movement within TME.



Patients with stroma-rich, fibrotic, and desmoplastic tumors such as pancreatic ductal adenocarcinoma (PDAC), and subsets of breast and ovarian adenocarcinomas or head-and-neck squamous carcinomas (HNSCC) face particularly poor outcomes.^{14–16} PDAC, in which the stromal compartment can constitute up to 90% of tumor volume, remains among the deadliest tumor entities despite significant therapeutic progress.^{17–19} Standard therapies and even modern immunotherapies often fail not because molecular targets are absent, but because dense stroma prevents drugs and immune cells from accessing tumor islands, creating immune-excluded or ‘cold’ tumors.^{20–26}

CAFs orchestrate this transport barrier through ECM deposition, fiber reorganization, and cytokine programs that spatially segregate immune cells and therapeutic agents.²⁷ For example, CXCL12 secretion sequesters T-cells in stromal compartments, a mechanism reversible by CXCR4 antagonists such as Plerixafor, revealing that matrix remodeling defines an immune transport dimension in tumors.²⁸ The TME therefore behaves as a dynamic and self-adapting transport system rather than a passive scaffold. CAFs, a heterogeneous population of tumor-reprogrammed stromal cells, continuously deposit and reorganize ECM components such as collagen and proteoglycans, establishing stiffness gradients and fiber architectures that sustain invasion and therapy resistance.^{18,21} Proliferation of cancer cells and CAFs generates excessive ECM and solid stress (SS) that compresses blood and lymphatic vessels, elevates interstitial fluid pressure (IFP), and disrupts perfusion, producing heterogeneous advective-diffusive transport regimes that limit drug penetration.^{29–33} Quantitative studies indicate that only a minute fraction of administered drugs reaches malignant cells, underscoring the dominant role of physical barriers.³³

Both cancer cells and CAFs sense and respond to these mechanical cues *via* mechanotransduction pathways (including integrins, the cytoskeleton, ion channels, and YAP/TAZ, MAPK signaling), converting forces into biochemical programs that further drive ECM remodeling and barrier reinforcement.^{34–40} This establishes a bidirectional mechanochemical feedback loop in which matrix mechanics and transport properties continuously co-evolve with stromal activation.^{38,41–45}

Consequently, each tumor develops a unique and evolving transport phenotype, defined by ECM architecture, intrinsic permeability, and interstitial flow patterns that govern the accessibility of drugs and immune cells. However, identifying and quantifying which features of this remodeled matrix impede therapeutic transport remains a major challenge. Microfluidics and tumor-on-chip (ToC) technologies have emerged to address this problem by combining physiological relevance with experimental control and transparency. Unlike 2D cultures or animal models, ToC platforms can reproduce stromal architecture, pressure gradients, oxygen and nutrient transport, and CAF-mediated ECM remodeling under defined conditions, while enabling direct measurement of stiffness, porosity, and molecular gradients.^{46–52} These systems therefore provide a unique opportunity to characterize patient-specific

transport phenotypes and to link CAF-driven matrix remodeling to therapeutic accessibility. The central motivation of this review is to examine how CAF-mediated mechanotransduction drives ECM remodeling and transport barrier formation, and how microfluidic and computational models can be used to quantify and perturb these transport phenotypes. Each approach provides distinct mechanistic insights and translational opportunities. We therefore structure this review around a causal and transport-centered framework. Section 2 establishes the biological foundations by defining the functional states of CAFs and their roles in ECM remodeling and mechanical feedback. Section 3 introduces the physical principles of mass transport in dynamically remodeling tissues and shows how these concepts can be experimentally accessed using microfluidic tumor-on-chip platforms. Section 4 integrates these biological and physical dimensions to analyze transport-limited, extrinsic mechanisms of therapeutic resistance driven by stromal remodeling. Finally, section 5 discusses how transport phenotyping can be leveraged as a functional complement to genomic profiling, with implications for patient stratification, therapeutic sequencing, and clinical translation.

2. The causality chain: a unifying framework for the impact of matrix remodeling on tumor biology

2.1 Conceptual foundations

The complexity of tumor–stroma interactions has historically resisted purely reductionist approaches. CAF heterogeneity, ECM composition, chemical gradients, and mechanical forces are coupled through nonlinear feedback loops that blur simple cause-and-effect relationships.^{10,11,53–55} This complexity calls for conceptual frameworks that integrate cellular diversity, matrix mechanics, and dynamic transport. CAF populations form a functional spectrum of overlapping states that jointly shape the TME.⁵⁶ Among these, we define matrix-producing and matrix-remodeling CAFs (mat/mrCAFs) as the CAF populations whose dominant functional program is ECM deposition, crosslinking, and fiber reorganization. This operational definition largely overlaps with the classical myofibroblastic CAF (myCAF) phenotype, characterized by high α -smooth muscle actin (α -SMA) expression and contractility, as well as strong expression of fibrillar collagens (COL1A1, COL3A1), fibronectin (FN1), and crosslinking enzymes such as LOX and LOXL1/2. These enzymes generate covalent collagen crosslinks that increase matrix stiffness while reducing porosity and intrinsic permeability.^{57,58} In contrast, inflammatory CAFs (iCAFs) display dominant cytokine and chemokine programs (*e.g.*, IL-6, CXCL12) that primarily modulate paracrine signaling, immune recruitment, and vascular function, while antigen-presenting CAFs (apCAFs) express MHC-II transcripts and participate in immune interactions.^{59–62} Although these CAF states overlap biologically and individual CAFs may display mixed

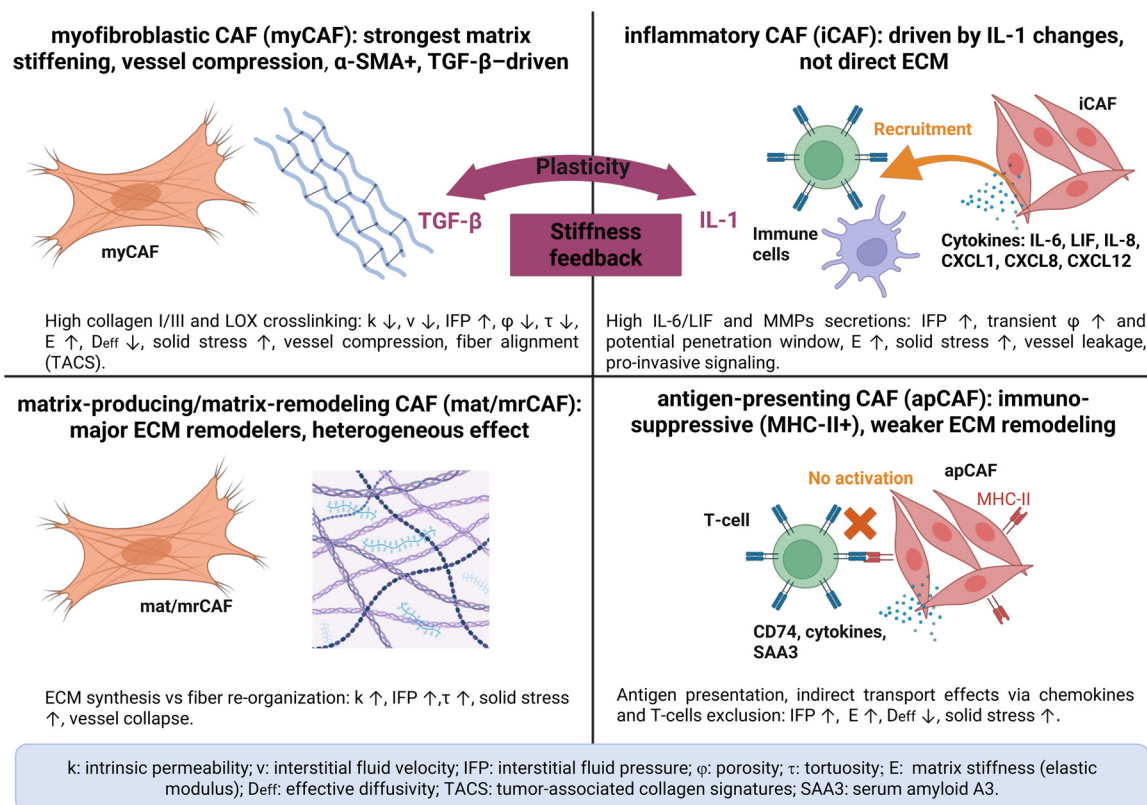


programs, their distinction is useful because their dominant functions have different consequences for tissue transport: mat/mrCAF matrix deposition reduces porosity and increases tortuosity; myCAF contractility generates solid stress and vessel compression; iCAF and apCAF programs indirectly modulate transport through the immune and vascular pathway. For clarity, we use the mat/mrCAF label as a shorthand when discussing matrix to transport coupling.

CAF heterogeneity extends beyond molecular markers to functional consequences on matrix architecture and transport properties, with distinct CAF states differentially shaping stiffness, porosity, and immune-cell accessibility (see Box 1).

These CAF populations are embedded in reciprocal mechanochemical feedback loops. Activated mat/mrCAFs deposit and reorganize ECM, increasing stiffness and generating fiber alignment and gel contraction. The remodeled matrix, in turn, alters the mechanical and biochemical environment sensed by CAFs, further reinforcing their activated state. Transport processes are an integral part of this loop: ECM remodeling modifies diffusivity, intrinsic permeability, and interstitial flow, thereby reshaping the spatial distributions of oxygen, nutrients, cytokines, immune cells, and therapeutic agents.⁶³ These altered gradients feedback on CAF activation, cancer cell plasticity, and immune

behavior, creating emergent transport barriers that are not fixed but dynamically maintained. Single-cell and functional studies illustrate how distinct CAF subtypes contribute to this process through complementary mechanical strategies. For example, CAF-S4/myCAF-like cells exhibit a highly contractile, pericyte-like program with elevated strain-energy density sufficient to compact local collagen and mechanically promote invasion, whereas CAF-S1/myCAF-like cells emphasize proliferative and secretory programs that further drive ECM production and cytokine-mediated crosstalk.⁶⁴ Together, these CAF subsets generate spatially organized mechanical and matrix architectures that define transport pathways within the tumor.^{65,66} At the tissue level, CAF-driven ECM deposition and contractility produce desmoplasia, increased solid stress, and vessel compression, leading to elevated interstitial fluid pressure and heterogeneous advective-diffusive transport regimes.⁶⁷ These physical changes translate directly into spatial gradients of oxygen, metabolites, drugs, and immune cells, which in turn govern cancer cell proliferation, migration, and survival. Histologically, this process manifests fibrotic stromal networks surrounding distorted tumor islands, abnormal vasculature, and infiltrative growth patterns that correlate with clinical aggressiveness and therapeutic resistance.^{65,66,68–75}



Box 1 Schematic overview of representative CAF functional states and their dominant effects on extracellular matrix (ECM) structure and mass transport in stroma-rich carcinomas

Upper left (myCAF): primary matrix-stiffening subtype; upper right (iCAF): inflammatory subtype with indirect transport effects *via* MMPs and vascular leakage; lower left (mat/mrCAF): heterogeneous ECM remodelers; lower right (apCAF): immunosuppressive subtype with indirect transport effects *via* T-cell



exclusion. The central element depicts bidirectional plasticity driven by TGF- β /IL-1 between subtypes. Myofibroblastic CAFs (myCAFs) are characterized by high α -smooth muscle actin (α -SMA) expression, contractility, and TGF- β -driven matrix crosslinking, leading to increased tissue stiffness, vessel compression, elevated interstitial fluid pressure, and reduced effective diffusivity and intrinsic permeability. Matrix-producing and matrix-remodeling CAFs (mat/mrCAFs) deposit, reorganize, and degrade ECM fibers, dynamically modulating porosity, tortuosity, and permeability, thereby generating spatially heterogeneous transport pathways. Inflammatory CAFs (iCAFs) secrete cytokines and chemokines (e.g., IL-6, CXCL12) that primarily affect immune-cell localization and vascular function, indirectly influencing transport. Antigen-presenting CAFs (apCAFs) express MHC class II molecules and modulate immunity, with secondary effects on immune infiltration and spatial drug distribution. Not all CAF states are present or dominant across all tumor types. The schematic illustrates typical functional programs reported in desmoplastic cancers and highlights how distinct CAF activities contribute, directly or indirectly, to the emergence of CAF-driven transport phenotypes that regulate drug and immune-cell accessibility.

Together, these observations motivate a causal framework in which CAF activation, ECM remodeling, and transport are tightly coupled (Fig. 1). Rather than representing linear cause-and-effect relationships, the arrows in this framework denote bidirectional feedback: CAFs remodel the matrix, the matrix alters transport, and transport reshapes cellular states. This reciprocal coupling can generate stable or evolving stromal-tumor configurations whose defining feature is a specific transport phenotype. Understanding and perturbing these CAF-driven transport phenotypes is therefore central to deciphering

therapy resistance and to designing effective tumor-on-chip experiments. Indeed, this framework naturally enables hypothesis-driven ‘virtual experiments’ by inverting inputs and outputs along the causality chain. Forward causality can be tested by perturbing upstream drivers of CAF activation and remodeling (e.g., TGF- β signaling or LOX activity) while tracking transport readouts, whereas reverse causality can be probed by directly modulating ECM-level variables (stiffness, porosity, intrinsic permeability, or interstitial pressure) and quantifying cellular responses. Microfluidic systems are

The causality chain framework for understanding matrix-transport coupling in tumors

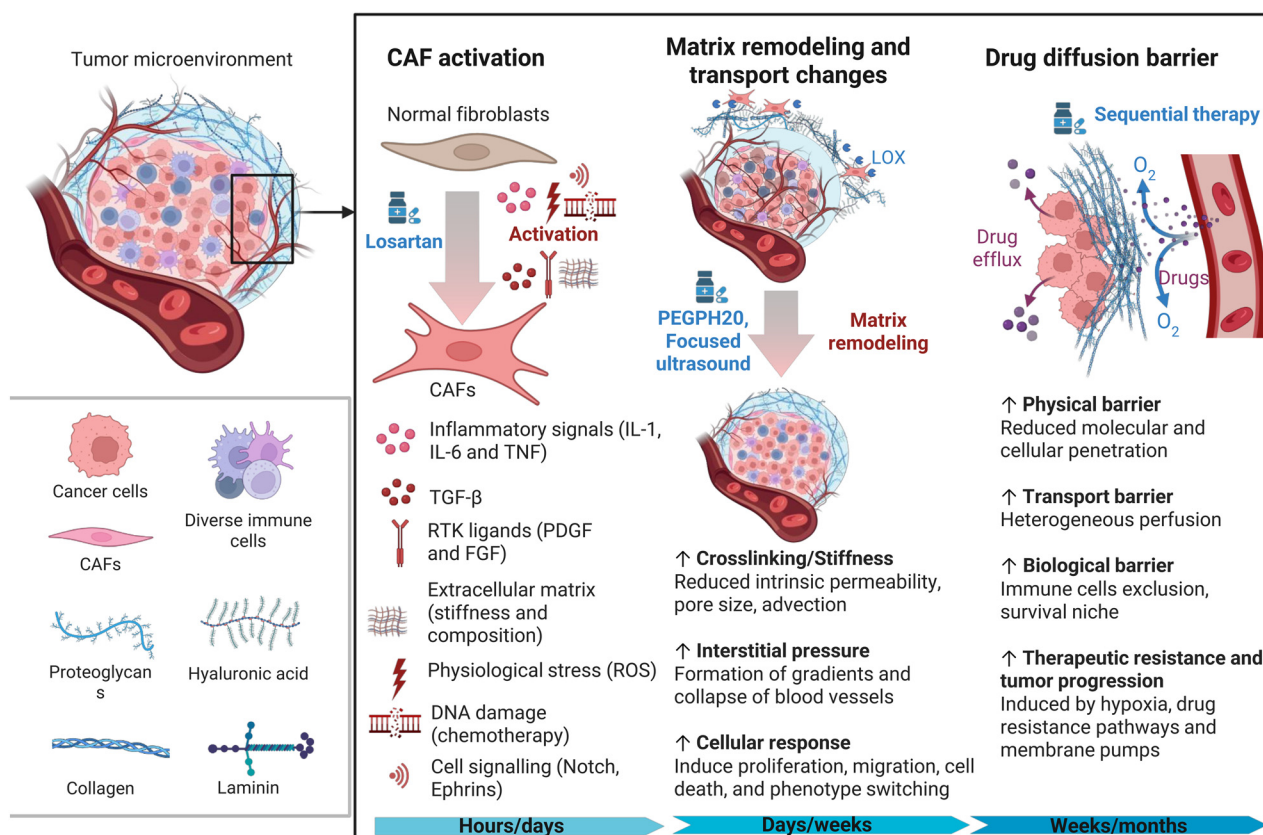


Fig. 1 Causality-chain schematic linking CAF activation, matrix/transport remodeling, and drug-diffusion barrier formation in desmoplastic tumors. A left-to-right sequential schematic depicts three representative, interdependent stages commonly observed in desmoplastic tumors and their reinforcing feedback: (1) CAF activation (hours–days) → (2) matrix remodeling & transport alteration (days–weeks) → (3) drug-diffusion barrier emergence (weeks–months). Stage summaries: (1) local biochemical and mechanical cues rapidly activate stromal fibroblasts; (2) activated CAFs remodel the extracellular matrix, producing structural and compositional changes that modify local diffusivity and advective transport; (3) accumulated remodeling yields persistent architectural barriers (compartmentalization, heterogeneous perfusion, and solid stress, immune exclusion) that impede drug and immune-cell access. Strategies are mapped (in blue) to stages to indicate where therapies could blunt activation, modify matrix structure, transiently enhance transport, or sequence treatments to exploit improved access. Temporal ranges are indicative and vary with tumor type and context.



particularly well suited for such bidirectional causal tests because they allow controlled tuning of boundary conditions and matrix properties together with real-time, multiparametric measurements, thereby providing a quantitative bridge between stromal mechanics and drug transport.^{76–87} When the ECM is considered as the primary causal input, relevant structural, mechanical, and molecular parameters, including stiffness, composition, porosity, permeability, and mechanotransduction readouts, can be quantified experimentally. A detailed list of input and output variables and their measurement modalities is provided in section S1.

2.2 Transport physics in dynamic biological systems

Eqn (1)–(7) together form a compact mathematical framework that links solute transport and tissue mechanics in CAF-remodeled tumors. Numerical and analytical solutions yield spatial and temporal maps of solute concentration, interstitial pressure, and tissue deformation, which directly reflect how ECM density, crosslinking, and solid stress shape drug and immune-cell accessibility. These maps address biologically central questions: where will a drug penetrate within a CAF-rich tissue? Is poor delivery dominated by limited diffusion through a dense collagen network, by advective wash-out driven by elevated interstitial fluid pressure, or by rapid cellular uptake? How will interventions that modulate CAF activity, alter matrix structure, or relieve solid stress reshape tissue exposure? For readers less familiar with mathematical notation, the equations can be read as a quantitative blueprint that connects CAF-driven ECM remodeling to measurable transport parameters such as effective diffusivity, intrinsic permeability, and interstitial pressure, and from there to spatial patterns of drug distribution that can be directly compared with microfluidic and tumor-on-chip experiments.^{12,13,88–92} A complete list of symbols, units, and their corresponding experimental readouts in tumor-on-chip systems is provided in Table S1.

Box 2 Mathematical framework for CAF-driven transport phenotyping

Understanding solute transport through tumor tissue requires applying physical principles to dynamic, evolving biological systems. In solid tumors, CAF activity, ECM remodeling, vascular leakiness, and cell contractility continuously reshape diffusion, advection, and tissue deformation. Unlike static engineered porous media, tumor transport therefore occurs in a matrix whose porosity, stiffness, and permeability evolve over time. The coupled advection–diffusion–reaction and poroelastic equations introduced in section 2.2 provide a compact but mechanistically meaningful way to formalize this complexity. Rather than being abstract, each variable appearing in the equations corresponds to a biologically and experimentally accessible property of CAF-remodeled tissue. For example, the effective diffusivity D_{eff} reflects how collagen density, fiber alignment, and crosslinking by mat/mrCAFs restrict molecular motion; the intrinsic permeability k captures how ECM compaction, LOX-mediated crosslinking, and cellular contraction regulate interstitial fluid flow; and the interstitial fluid pressure p reports the combined effects of vascular leakiness and CAF-driven solid stress.

In this formulation, matrix remodeling and mechanotransduction directly enter the transport problem through their impact on these coefficients. Increased matrix deposition and crosslinking by mat/mrCAFs reduce ϕ , increase tortuosity, and lower both D_{eff} and k , while myCAF contractility elevates solid stress and interstitial pressure, thereby modifying advective transport. Conversely, enzymatic matrix degradation, mechanical decompression, or vascular normalization act by shifting the same parameters in the opposite direction. The mathematical framework therefore translates qualitative descriptors such as ‘dense’, ‘stiff’, or ‘fibrotic’ into quantitative variables that can be measured, quantitative variables that can be measured, perturbed, and used to test biological hypotheses. By linking CAF activity and ECM structure to transport coefficients that determine drug and immune-cell access, this formalism provides a quantitative bridge between stromal biology and therapeutic delivery and defines the basis for transport phenotyping in tumor-on-chip systems.

For readers less familiar with mathematical formalism, the conceptual role of the equations introduced below and their relationship to experimentally measurable biological quantities are summarized in Box 2.

2.2.1 Solute/mass balance: advection–diffusion–reaction.

The generalized advection–diffusion–reaction equation describes how diffusion, interstitial flow, and cellular uptake together determine the spatiotemporal distribution of therapeutic molecules in tumor tissue.^{34,90,93}

$$\frac{\partial c}{\partial t} = \nabla \cdot (D_{\text{eff}}(\phi, x, y, z, t) \nabla c) - \nabla \cdot (v_p c) + R(x, y, z, t) \quad (1)$$

Here, c is the local solute concentration, D_{eff} is the effective diffusivity in the CAF-remodeled extracellular matrix, v_p is the interstitial fluid velocity, and R represents cellular uptake, degradation, or local secretion.

In CAF-rich tumors, none of these terms are spatially or temporally constant.^{94,95} The effective diffusivity D_{eff} reflects how matrix-producing and matrix-remodeling CAFs alter collagen density, fiber alignment, and crosslinking. Increased deposition of fibrillar collagens and LOX-mediated crosslinking reduce matrix porosity ϕ and increase tortuosity λ , thereby lowering $D_{\text{eff}} = D_0 \phi / \lambda^2$ relative to free diffusion in the interstitial fluid.^{96–99} As a result, even small molecules experience slowed penetration in desmoplastic regions, while large biologics or nanoparticles are much more strongly hindered.

The advective term $\nabla \cdot (v_p c)$ captures the contribution of interstitial fluid flow to solute transport. In tumors, CAF-driven matrix contraction, vessel compression, and vascular leakiness generate elevated and spatially heterogeneous interstitial fluid pressure, producing local fluid velocities that can either enhance delivery along pressure gradients or promote convective wash-out from poorly perfused tumor regions. Thus, CAF activity enters the transport equation not only through diffusion but also through pressure-driven advection.



The reaction term $R(x, y, z, t)$ accounts for cellular uptake, binding, or degradation of the solute. In addition to cancer-cell-intrinsic drug consumption, CAFs themselves can sequester, metabolize, or exclude therapeutic agents through ECM binding and pericellular matrix deposition. Consequently, the apparent rate of drug loss within a CAF-rich microenvironment reflects both cellular and matrix-associated sinks.

Unlike engineered porous materials, tumor tissues evolve continuously: CAF activation, ECM remodeling, vascular collapse or normalization, and cellular turnover all modify D_{eff} , v_p , and R over time. The advection–diffusion–reaction equation therefore provides a dynamic, mechanistically grounded description of how CAF-driven matrix remodeling gives rise to heterogeneous and time-dependent transport phenotypes that govern drug accessibility.^{97,98}

2.2.2 Interstitial fluid flow: Darcy-type transport. To define the advective term in eqn (1), interstitial fluid motion through the CAF-remodeled extracellular matrix is described by a Darcy-type law for porous media:

$$q = -k(\phi, x, y, z, t)/\mu \cdot \nabla p \quad (2)$$

where q is the Darcy volumetric flux, k is the intrinsic permeability of the matrix, μ is the dynamic viscosity of the interstitial fluid and p is the interstitial fluid pressure (IFP). In tumors, IFP is elevated by vascular leakiness, impaired lymphatic drainage, and CAF-driven solid stress that compresses blood and lymphatic vessels. Intrinsic permeability k captures how easily fluid can flow through the ECM and is therefore a direct physical readout of matrix architecture. In CAF-rich tissues, collagen deposition, fiber alignment, and LOX-mediated crosslinking reduce pore connectivity and increase tortuosity, leading to large decreases in k . Similarly, myCAF contractility compacts the matrix and mechanically constricts interstitial spaces, further lowering permeability. Conversely, enzymatic matrix degradation or mechanical decompression transiently increases k by reopening fluid pathways.

The interstitial fluid velocity v_p entering eqn (1) is described by:

$$v_p = q/\phi(x, y, z, t) = -(1/\phi)k(\phi, x, y, z, t) \cdot \nabla p. \quad (3)$$

Darcy's law is analogous to water percolating through a sponge: fluid moves faster when permeability is higher, viscosity is lower, or pressure gradients are steeper. This expression makes explicit how CAF-driven changes in porosity ϕ , intrinsic permeability k , and pressure gradients together determine advective transport. In desmoplastic tumors such as PDAC, low permeability combined with high IFP produces steep pressure gradients that can promote convective wash-out of drugs from tumor cores and hinder inward penetration from blood vessels.^{100–102}

Darcy's law thus provides a mechanistic link between stromal remodeling and drug delivery: CAF-mediated matrix

densification and vessel compression simultaneously reduce permeability and elevate pressure, shifting transport from diffusion-dominated regimes toward outward, pressure-driven flow. These coupled effects are central to the emergence of CAF-driven transport barriers that limit therapeutic and immune-cell access.

2.2.3 Solid deformation: poro(visco)elastic mechanics. While eqn (1) captures solute balance and eqn (2) describes fluid flow, neither accounts for the fact that the tumor ECM is a deformable, cell-driven material. In CAF-rich tumors, matrix architecture is continuously reshaped by cell-generated forces, which in turn modulate porosity, permeability, and diffusivity.⁹³ This mechanical coupling is captured by a poro-(visco)elastic description of the tissue.^{92,103} Mechanical equilibrium, the balance of forces within the solid matrix is expressed as:

$$\nabla \cdot \sigma + b = 0, \quad (4)$$

where σ is the stress tensor and b represents body forces. In biological terms, these stresses arise from myCAF contractility, proliferating cancer cells, and externally imposed compression from the surrounding tissue.

To relate stress to deformation, poroelastic or poroviscoelastic constitutive laws are used. In this framework, tissue stress depends both on solid deformation and on pore pressure.^{103,104} A Kelvin–Voigt-type formulation,

$$\sigma = E \cdot \varepsilon(u) + \eta \frac{\partial \varepsilon(u)}{\partial t} \alpha p \quad (5)$$

links the elastic stiffness E and viscous relaxation η of the ECM to interstitial fluid pressure p , with α describing how pore pressure contributes to solid stress.

In CAF-remodeled tissues, these coefficients are not material constants but biological variables. Matrix deposition and LOX-mediated crosslinking by mat/mrCAF increase the effective elastic modulus E , while myCAF contractility and cell proliferation generate compressive strain ε that compacts the matrix. Elevated pore pressure p , resulting from vascular leakiness and vessel compression, further contributes to solid stress. Together, these effects lead to progressive densification of the ECM, reduction of pore volume, and distortion of fluid pathways.

Through this coupling, mechanical remodeling feeds directly back into transport.^{92,95} Local increases in stiffness and compressive strain reduce porosity ϕ and intrinsic permeability k , while elevated pore pressure modifies advective flow. The poro-(visco)elastic formulation therefore provides a mechanistic description of how CAF-driven force generation and matrix stiffening translate into dynamic, spatially heterogeneous transport barriers within solid tumors.^{105,106}

2.2.4 Fluid mass conservation and solute storage. The poroelastic formulation above integrates solid deformation and fluid pressure, but transport also depends on how much



fluid and solute a deforming matrix can store. This coupling is captured by the fluid mass conservation equation:

$$\nabla \cdot q + \partial/\partial t[\alpha(\nabla \cdot u) + (1/M)p] = Q_f \quad (6)$$

where q is the interstitial fluid flux, p is the interstitial fluid pressure, M is the Biot storage modulus, α couples solid deformation to fluid content, and Q_f represents vascular leakage or drainage.

In CAF-rich tumors, this equation formalizes how matrix compaction and vessel compression alter fluid storage. MyCAF-driven contraction and proliferative growth of tumor cells reduce pore volume, expelling fluid and raising interstitial pressure, whereas vascular leakiness continuously injects fluid into the tissue. The balance of these processes determines whether interstitial flow stagnates or whether pressure-driven fluid motion dominates.

Because porosity changes with deformation, solute storage must evolve accordingly. A generalized solute mass balance including sorption is:

$$\begin{aligned} \partial/\partial t[\phi c + (1 - \phi)c_s] + \nabla \cdot (cq - \phi D_{\text{eff}}(\phi, x, y, z, t) \cdot \nabla c) \\ = R(c, p, u, x, t) + Q_c \end{aligned} \quad (7)$$

Here, ϕ is the local porosity, c the mobile solute concentration, and c_s the fraction bound or immobilized within the solid matrix (for example through binding to collagen, proteoglycans, or pericellular ECM deposited by CAFs). Q_c represents local solute sources or sinks.

CAF-driven ECM deposition, crosslinking, and compression decrease ϕ , reducing the volume available for fluid and solute transport while increasing the fraction of drug that becomes trapped or immobilized within the dense matrix. Conversely, enzymatic matrix degradation or mechanical decompression increases pore volume and reduces solute sequestration, transiently enhancing drug penetration and retention.

This formulation therefore captures an important biological reality: CAF-mediated matrix remodeling does not only alter how fast drugs move, but also how much of a drug can be stored, retained, or excluded within the tumor microenvironment, contributing to heterogeneous and time-dependent therapeutic exposure.^{93,95,97}

2.2.5 Experimental calibration of model parameters. Eqn (1)–(7) provide a mechanistic framework for coupled solute transport and tissue mechanics in CAF-remodeled tumors. However, their predictive value depends critically on experimental calibration. Key biophysical quantities, such as effective diffusivity $D_{\text{eff}}(\phi, x, y, z, t)$, intrinsic permeability $k(\phi, x, y, z, t)$, reaction and uptake rates $R(x, y, z, t)$, and poroelastic coefficients such as E , η , α , and M , must be measured in biologically relevant stromal environments. Recent advances in microfabricated tumor–stroma models and tissue-mimetic hydrogels make such calibration feasible.^{80,107–109} These systems allow controlled variation of matrix density, crosslinking, CAF composition, and

interstitial pressure while providing optical and sensor access to transport and mechanical readouts. In this context, CAF-driven ECM remodeling becomes an experimentally tunable input, and transport coefficients become measurable outputs.

For example, collagen content and LOX-mediated crosslinking determine matrix stiffness and pore architecture, which can be quantified by rheology, AFM, and imaging-based measurements of porosity and fiber organization.^{77,78,82,110,111} These structural properties directly constrain D_{eff} and k , which can be extracted from tracer diffusion, FRAP, or pressure-driven flow assays in microfluidic devices.^{78,82} Similarly, myCAF contractility and tissue compression modulate solid stress and interstitial pressure, which can be measured through pressure sensors, deformation tracking, or poroelastic relaxation experiments.^{84,112,113}

By fitting the transport and poroelastic equations to experimental data, one can infer spatially and temporally resolved transport phenotypes that reflect the underlying CAF and ECM state. Importantly, this calibration is not limited to static conditions: microfluidic systems allow perturbation of CAF activity (e.g., TGF- β inhibition, LOX blockade), matrix structure (enzymatic degradation, crosslinking), or mechanical loading, enabling direct tests of how specific stromal interventions reshape D_{eff} , k , and p , and thereby drug accessibility.

In this way, tumor-on-chip platforms transform the mathematical framework into a practical tool for CAF-driven transport phenotyping, linking stromal biology, physical transport parameters, and therapeutic response in a quantitatively testable manner.

2.3 Designing experiments to extract CAF-driven transport phenotypes

The central challenge in tumor transport modeling is not mathematical formulation but experimental calibration: the deliberate design of experiments that yield biologically meaningful values for effective diffusivity D_{eff} , intrinsic permeability k , and reaction terms R in CAF-remodeled tissues. Because these parameters emerge from the cellular and stromal architecture of tumors, experimental systems must capture both matrix mechanics and CAF-driven remodeling.^{12,13,34,114,115}

Microfluidic tumor–stroma platforms provide a uniquely suited environment for this task. By controlling CAF composition, ECM density, and interstitial pressure while continuously measuring tracer transport and tissue deformation, these systems enable direct mapping of stromal state to transport phenotype. In practice, this means that CAF populations (for example, myCAF-enriched *versus* mat/mrCAF-enriched mixtures) can be embedded in collagenous matrices, exposed to defined mechanical or biochemical perturbations, and probed using fluorescent tracers or therapeutic molecules to quantify penetration and retention



over time. A practical entry point is to extract spatially averaged transport coefficients by fitting advection–diffusion–reaction models to tracer uptake or release data. For example, time-resolved penetration of fluorescent dextran derivatives, albumin, or chemotherapeutics in CAF-conditioned matrices can be fitted to infer D_{eff} and apparent uptake rates.^{107–109,116}

Similarly, pressure-relaxation or flow-through assays yield estimates of intrinsic permeability and elastic moduli, directly reporting how CAF-driven matrix densification and contraction modulate interstitial transport.^{117–119}

Importantly, CAF-mediated remodeling is inherently heterogeneous and dynamic. As CAFs deposit, crosslink, and reorganize collagen, local values of D_{eff} and k evolve over days to weeks. Microfluidic platforms allow these temporal changes to be tracked and, when combined with imaging of collagen architecture or CAF localization, provide spatially resolved transport phenotypes that reflect the underlying stromal organization.

Detailed descriptions of hydrogel systems, tracer-based diffusion measurements, pressure–flow assays, and sources of experimental variability, as well as their use for quantitative model calibration, are provided in section S2, which presents the full methodological and materials-science framework supporting this approach.

By systematically perturbing CAF activity (e.g., TGF- β blockade, LOX inhibition), matrix structure (enzymatic degradation, crosslinking), or mechanical loading (compression, stress relief), one can establish causal links between stromal biology and transport.^{120,121} In this way, tumor-on-chip systems enable bidirectional testing: CAF-driven matrix remodeling predicts transport behavior, and measured transport phenotypes, in turn, reveal the dominant stromal mechanisms that limit drug delivery. These CAF-driven transport parameters do not change monotonically but evolve through characteristic temporal phases during tumor progression, yielding transient windows of enhanced accessibility, followed by the establishment of persistent transport barriers. A conceptual synthesis of these dynamics, integrating experimental observations and tumor-on-chip studies, is illustrated in Fig. 2.

2.4 Clinical implications: from CAF-driven transport phenotypes to therapeutic stratification

Modular *in vitro* tumor–stroma models provide a practical route to translate CAF-driven transport biology into clinically actionable information. By explicitly manipulating fibroblast populations, ECM composition, and mechanical loading, these systems allow direct measurements of how stromal remodeling shapes drug and immune accessibility.

Patient-derived CAFs or defined CAF subtypes can be combined with controlled ECM formulations to generate tissues with distinct transport phenotypes, characterized by their effective diffusivity, intrinsic permeability, and interstitial pressure.^{122,123} Fluorescent tracers or therapeutic

agents introduced into these systems reveal how rapidly and how deeply molecules penetrate, providing a functional readout of stromal barrier strength. When integrated with transport models, these measurements identify transient windows of enhanced accessibility or persistent regions of exclusion. This approach has immediate clinical implications. Tumors that appear similar at the genomic or histological level can differ markedly in their CAF composition and matrix architecture, leading to very different transport phenotypes. Such differences help explain why drugs that are highly potent *in vitro* or in genetically stratified patients fail to achieve sufficient exposure in desmoplastic tumors such as PDAC, breast, or ovarian carcinomas.^{13,29,124,125} Transport phenotyping therefore provides a complementary axis of patient stratification, orthogonal to molecular profiling.

Importantly, CAF-driven transport barriers are not fixed. Enzymatic matrix remodeling, mechanical stress relief, or pharmacological reprogramming of CAFs can transiently increase permeability and diffusivity, thereby improving drug penetration. Tumor-on-chip platforms allow these interventions to be tested in a controlled and patient-specific manner, enabling rational sequencing of stromal-modulating and cytotoxic or immunotherapeutic agents.^{125–127}

A detailed overview of experimental and pharmacological strategies to modulate CAF-driven matrix barriers, including enzymatic, mechanical, and stromal reprogramming approaches, is provided in section S3.

By linking CAF biology to measurable transport parameters, and by providing experimental systems to perturb both, transport phenotyping offers a framework to identify which patients are likely to benefit from stromal targeting, and when such interventions should be applied to maximize therapeutic access.

3. Microfluidic platforms: bridging scales and transport complexity

3.1 From model hierarchy to transport-resolved tumor-on-chip systems

The evolution from conventional *in vitro* tumor models to microfluidic ToC platforms has been driven by a central limitation of standard systems: their inability to resolve spatially heterogeneous transport under dynamic stromal remodeling. While 2D monolayers, spheroids, and organoids reproduce selected aspects of tumor organization, they fail to capture the coupled gradients of nutrients, oxygen, drugs, and mechanical stresses that emerge in confined, perfused, and remodeling matrices.¹²⁸ Microfluidic ToC systems uniquely address this gap by combining controlled geometry, perfusion, and mechanical confinement with three-dimensional cell–matrix organization. These platforms enable the direct measurement of interstitial flow, pressure gradients, and solute transport within evolving stromal architectures, thereby linking CAF activity, matrix remodeling, and drug accessibility in a single experimental



Temporal evolution of transport properties during tumor progression

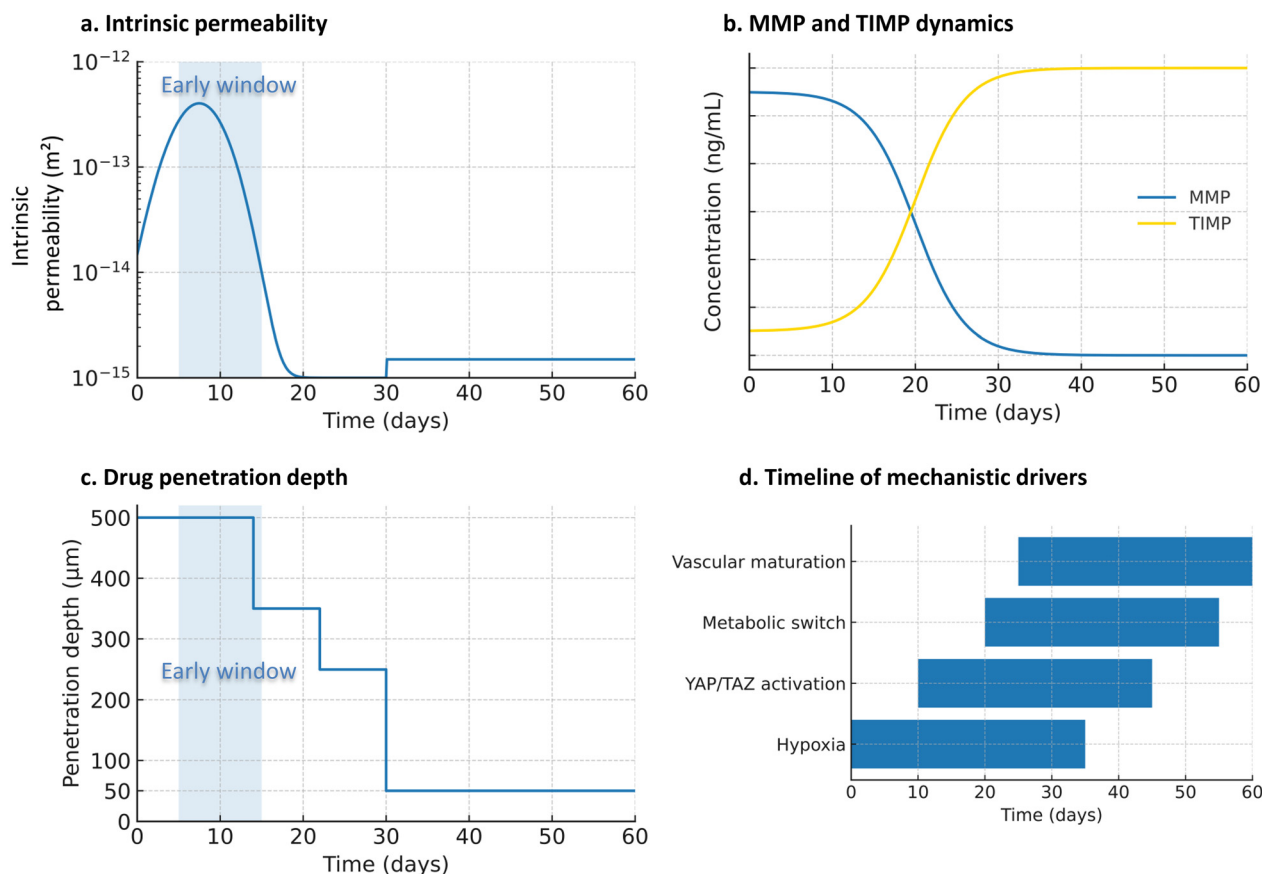


Fig. 2 Conceptual timeline of CAF-driven transport remodeling during tumor progression. This figure presents a conceptual, hypothesis-driven schematic summarizing recurrent trends reported across experimental, preclinical, and microfluidic tumor-on-chip studies, rather than quantitative data derived from a single system. The timelines illustrate how CAF-mediated matrix remodeling progressively alters transport properties during tumor evolution, with indicative temporal scales that vary depending on tumor type, stromal composition, and experimental context. **a.** Schematic evolution of intrinsic permeability highlighting an early permissive window characterized by relatively high permeability, followed by a transition phase and the establishment of a transport-limiting stromal barrier associated with matrix densification and solid stress accumulation (typical ECM range: 10^{-15} – 10^{-13} m^2). **b.** Qualitative dynamics of matrix metalloproteinase (MMP) activity and tissue inhibitors of metalloproteinases (TIMPs), illustrating an early remodeling-dominated phase (MMP-biased) followed by progressive inhibition and matrix stabilization in later stages. **c.** Representative evolution of drug penetration depth, illustrating deep tissue access during early stages and progressive exclusion as matrix density, tortuosity, and interstitial pressure increase. **d.** Indicative timing of major mechanistic drivers frequently associated with stromal maturation, including hypoxia, YAP/TAZ mechanotransduction, metabolic reprogramming, and vascular remodeling. Blue shading highlights potential therapeutic windows during which transient modulation of stromal or mechanical properties may improve drug accessibility.

framework.¹²⁹ In desmoplastic tumors, where CAF-driven matrix densification and solid stress dominate transport resistance, this capability is essential.^{71–73,130,131} An overview of the hierarchy of tumor models used for mass transport studies, including sealed and openable tumor-on-chip configurations, is provided in Fig. S1. Material-dependent transport artefacts and their implications for quantitative transport assays are summarized in Table S2.

3.2 Core microfluidic architectures for transport phenotyping

Despite wide design diversity, most transport-focused ToC platforms fall into three functional classes, defined by their ability to impose and measure transport-relevant fields.

(1) High-throughput spheroid generators. Droplet- or jet-based microfluidic systems enable rapid production of uniform spheroids or cell-laden microgels and are well suited for screening cytotoxic responses. However, because they lack controlled interstitial advection and matrix-scale pressure gradients, they are poorly suited for mechanistic studies of CAF-driven transport heterogeneity.^{52,132,133}

(2) Compartmentalized perfused chips with central gel regions. Perfused chips containing a confined central gel region flanked by flow channels constitute the workhorse architecture for transport studies. In these systems, CAFs and tumor cells remodel a three-dimensional ECM while defined pressure or flow differences generate interstitial advection. This configuration enables quantitative measurements of



permeability tracer washout, solute penetration depth and spatial transport heterogeneity as matrix density and stiffness evolve. Numerous studies across breast, colorectal, and pancreatic cancer models demonstrate that CAF-mediated matrix remodeling progressively reduces effective diffusivity and intrinsic permeability, directly limiting drug access.^{51,134}

(3) Integrated and openable platforms for multimodal characterization. More advanced ToC designs incorporate direct access to the matrix compartment or integrate *in situ* mechanical and biochemical measurements. Openable chips and multilayer platforms allow stiffness mapping, collagen imaging, and metabolic profiling to be performed on the same tissue that undergoes transport measurements.¹³⁵ These systems provide a unique opportunity to correlate ECM architecture, poroelastic properties, and drug penetration within a single experimental timeline, although real-time integration of all readouts remains technically challenging.^{135–140}

Material-specific transport artefacts and surface functionalization strategies are discussed in section S4.

3.3 Toward longitudinal, transport-resolved tumor models

A recurring limitation of current ToC platforms is their predominantly static or endpoint characterization of transport. CAF-driven remodeling, however, is inherently dynamic: intrinsic permeability, porosity, and solid stress evolve over days to weeks as matrix deposition, contraction, and reorganization progress.^{9,10,115,141–144} Capturing these dynamics, therefore requires longitudinal platforms capable of repeated or continuous measurements of key transport parameters, including intrinsic permeability, interstitial velocity, and solute diffusion.^{51,52,86,107–109,116}

Because transport properties are inseparably linked to the physicochemical and mechanical characteristics of the surrounding extracellular matrix, the choice of hydrogel and ECM formulation becomes a central experimental determinant rather than a neutral scaffold. Different matrices offer distinct ranges of stiffness, degradability, and pore architecture, which directly condition the accessible transport regimes and the interpretation of CAF-driven remodeling effects. The implications of hydrogel and ECM choice for CAF-driven transport phenotyping, including tunability of transport parameters and associated experimental pitfalls, are summarized in Table S3.

However, dynamic CAF-driven remodeling not only alters transport coefficients; it can also induce matrix contraction, densification, and geometric distortion of the gel compartment itself.^{48,145,146} If not adequately controlled, these effects confound transport measurements by modifying flow paths, pressure gradients, and effective diffusion distances over time. Consequently, reliable longitudinal transport phenotyping requires actively remodeling matrices to be mechanically and chemically stabilized within the microfluidic device. Detailed hydrogel

stabilization and anchoring strategies are described in sections S5 and S6.

Future ToC systems should therefore prioritize (i) stable confinement of actively remodeling matrices, (ii) reproducible control of pressure and flow boundary conditions, and (iii) integration of transport assays with structural and mechanical characterization. When combined with continuum or poroelastic transport models, such platforms enable the extraction of physically meaningful parameters that define a tumor's transport phenotype rather than a single drug response.^{147,148}

3.4 Technical limitations and methodological caveats

Despite their increasing sophistication, tumor-on-chip platforms remain constrained by methodological and technical limitations that must be considered when interpreting transport measurements. No single experimental technique simultaneously captures solute transport, matrix structure, and mechanics without bias. Fluorescent-based methods such as FRAP and fluorescent correlation spectroscopy (FCS) enable estimation of effective diffusivity but can locally perturb the system through photobleaching or photochemical damage and are limited to a small spatial sampling volume.^{116,149} Single-particle tracking (SPT) approaches often rely on nanometer-scale tracers (20–100 nm) that exceed the effective pore size relevant for small-molecule drugs, complicating quantitative extrapolation.¹⁵⁰

Structural imaging techniques, including second-harmonic generation (SHG) and confocal reflectance microscopy, provide detailed information on collagen organization and fiber alignment but do not directly report transport parameters.^{151,152} Mechanical mapping approaches such as atomic force microscopy (AFM), Brillouin microscopy, and elastography probe distinct mechanical regimes and length scales, each with intrinsic limitations in depth penetration, spatial resolution, or biological interpretation.¹⁵³ Moreover, labeling strategies can alter tracer size, charge, or hydrophobicity, thereby modifying transport behavior relative to unlabeled compounds.¹⁵⁴ These constraints underscore the necessity of multimodal experimental strategies that combine direct transport assays, structural and mechanical characterization, and mechanistic modeling. Cross-validation across techniques is essential to disentangle biological effects from methodological artefacts and to derive physically meaningful estimates of effective diffusivity, intrinsic permeability and poroelastic properties in evolving tumor microenvironments. Detailed methodological considerations and reporting recommendations are provided in section S7. Beyond these technical aspects, several unresolved conceptual and experimental challenges currently limit the quantitative interpretation and comparison of transport measurements across tumor-on-chip platforms; these are summarized in Box 3.



Box 3 Open questions on transport and mechanical confinement in tumor-on-chip systems

This Box highlights unresolved conceptual and experimental challenges that currently limit the quantitative interpretation of transport phenotypes in tumor-on-chip systems, particularly in the context of CAF-driven matrix remodeling and mechanical confinement.

1. Dynamic evolution of transport parameters under CAF-driven remodeling

How do key transport parameters, such as intrinsic permeability ($k(\phi, x, y, z, t)$) and effective diffusivity ($D_{\text{eff}}(\phi, x, y, z, t)$), evolve over time as CAFs remodel the extracellular matrix and generate solid stress? How can these dynamics be captured without assuming static tissue properties, given the continuous and nonlinear nature of stromal remodeling?

2. Separating device-induced artifacts from biological transport barriers

How can transport limitations arising from device-related effects, such as material sorption, gas permeability, or surface chemistry, be disentangled from genuine biological barriers created by CAF-driven ECM densification, fiber alignment, and stress-induced confinement?

3. Independent and coupled roles of solid stress and interstitial pressure

How can solid stress and interstitial fluid pressure be independently imposed or to identify their respective and combined contributions to drug, oxygen, and immune-cell transport within confined tumor microenvironments?

4. Coupling vascular dysfunction with interstitial transport

How can perfusion, endothelial permeability, and impaired lymphatic drainage be integrated with stromal remodeling to reproduce hypoperfusion-high IFP regimes characteristic of desmoplastic tumors?

5. Immune-cell transport in mechanically confined, CAF-remodeled matrices

How do chemotaxis, infiltration, and persistence of immune cells change as CAF-driven matrix density, fiber alignment, and compressive stress are systematically varied? To what extent does mechanical confinement contribute to immune exclusion independently of immunosuppressive signaling?

6. Toward comparable transport phenotypes across platforms

Which minimal set of transport- and mechanics-related parameters, such as ECM stiffness, intrinsic permeability, effective diffusivity, pressure gradients, porosity, fiber anisotropy, or confinement indices, is required to enable meaningful comparison and interpretation of transport phenotypes across tumor-on-chip studies?

3.5 Patient-derived systems and translational relevance

Patient-derived ToC platforms extend transport phenotyping beyond generic models by preserving native stromal composition, CAF subtype balance, and ECM architecture. Unlike genomic profiling, which identifies molecular

vulnerabilities, these systems directly probe whether therapeutic agents can physically reach malignant cells within their microenvironment.

Experimental evidence increasingly shows that tumors with higher effective permeability or lower matrix-induced resistance exhibit faster and stronger responses to therapy, whereas dense, CAF-rich matrices delay drug accumulation and blunt efficacy.¹⁵⁵ By quantifying transport parameters under physiologically relevant flow conditions, patient-derived ToC models provide a functional complement to molecular biomarkers and offer a framework to stratify tumors based on accessibility rather than target expression alone.

Despite encouraging validation studies, clinical translation remains limited by assay duration, technical complexity, and regulatory constraints. Nevertheless, transport-focused ToC systems represent a promising intermediate scale between mechanistic modeling and clinical pharmacokinetics, particularly for desmoplastic tumors where physical barriers dominate therapeutic failure.

Given the strong sensitivity of transport measurements to experimental conditions and inter-laboratory variability, recommended minimum reporting standards and reproducibility considerations for tumor-on-chip transport studies are summarized in section S8.

The ability of tumor-on-chip platforms to quantify evolving transport barriers provides the experimental foundation for transport phenotyping. In the following section, we examine how these platforms are used to interrogate CAF-driven matrix remodeling as a modifiable therapeutic target.

The value of experimental tumor models lies less in predicting clinical outcomes with absolute precision, a goal unattainable given the multifactorial nature of efficacy, toxicity, and clearance, but rather in identifying the key biophysical determinants of drug accessibility.

Microfluidic systems incorporating patient-derived components show exceptional promise in this regard.^{156,157}

Beyond the use of primary patient cells, their advantage lies in retaining and reproducing native heterogeneity, including the relative abundance of CAF subtypes (mat/mrCAFs, iCAFs, pCAFs) and the ECM properties they generate, such as composition, stiffness, and spatial organization. Because both cellular composition and ECM architecture regulate interstitial transport, patient-derived ToC platforms offer a practical means to recreate clinically relevant barriers to drug penetration and chemosensitivity. Indeed, CAF heterogeneity produces stromal microenvironments that differentially modulate tumor growth and therapy response,¹⁵⁸ while increased collagen crosslinking elevates solid stress and reduces pore size, thus limiting drug delivery. By maintaining controlled tumor–stroma ratios and ECM features, these platforms enable the systematic analysis of how CAF subtype balance matrix density and crosslinking may influence drug distribution and efficacy.^{159–161}



Perfused microfluidic tumor platforms using primary patient-derived tissue now enable quantitative mapping of intratumoral transport under physiologically relevant flow conditions. For instance, interstitial fluid velocities of $\sim 0.1\text{--}10\ \mu\text{m s}^{-1}$, comparable to those measured *in vivo*, have been successfully reproduced in 3D ToC models, eliciting similar transport responses.¹⁶²

While genomic biomarkers identify molecular susceptibilities within the tumor cells, they provide no information about how drugs may reach those cells within their physical microenvironment. Patient-derived microfluidic systems complement these molecular assays by offering a functional transport-based phenotype. They permit direct *ex vivo* testing of chemosensitivity and reveal how tissue architecture, stromal density, vascular leakiness, and perfusion collectively govern drug distribution and efficacy.^{95,163,164} Tumors exhibiting higher permeability respond more rapidly and strongly to treatment. In contrast, dense and transport-limiting matrices delay drug accumulation and require more prolonged exposure to achieve comparable therapeutic effects.

Clinical and preclinical validation studies have shown that ToC platforms can reproduce therapeutic responses with remarkable fidelity across biological scales. In an esophageal adenocarcinoma model, neoadjuvant chemotherapy responses were accurately predicted for all eight patients within 12 days, matching the parallel clinical results.¹⁶⁵ Comparable results were obtained in breast and prostate tumor slice cultures, which reproduced the *in vivo* sensitivity to cisplatin and apalutamide, respectively.¹⁶⁶ Similarly, pancreatic cancer tumor chips based on hydrogel microcapsules generated patient-specific drug-response profiles consistent with clinical follow-up data.¹⁶⁷ Preclinical models reinforce these findings. Petreus *et al.* (2021) reported almost identical irinotecan efficacy between microfluidic tumor spheroids (52% growth inhibition) and xenograft (53%), confirming quantitative alignment between these two very different platforms.¹⁶⁸ Similarly, Toley *et al.* (2013) showed that reproducing doxorubicin release from liposomal formulations (Doxil) in a tumor-mimetic chip required explicit modeling of diffusion, binding, and clearance to match clinical outcomes.¹⁶⁹

Collectively, these studies illustrate how patient-derived and preclinical ToC platforms can bridge the gap between mechanistic experimentation and clinical pharmacokinetics, complementing molecular biomarker data by providing direct, physical insight into drug accessibility within the TME. Despite these advances, substantial barriers still limit the clinical integration of ToC systems. To date, no ToC platform has received formal FDA clearance or CLIA certification for clinical use with patients (referring to the Clinical Laboratory Improvement Amendments, the U.S. regulatory framework that governs the quality standards and accreditation of clinical laboratories authorized to perform diagnostic testing on human specimens). Although both the FDA and CDC are actively evaluating these technologies

through pilot initiatives such as the IStand program (Innovative Science and Technology Approaches for New Drugs (ISTAND) Program|FDA), broader regulatory adoption remains constrained by challenges in standardization, validation, and the definition of clear contexts of use.^{170,171}

Beyond the regulatory dimension, practical barriers persist. Assay turnaround time remains a significant limitation: while microfluidic drug transport assays based on established cell lines can provide readouts within 5–7 days,¹⁶⁸ workflows involving patient-derived material often require 2–4 weeks, or even up to 6–8 weeks, when patient-derived organoids (PDOs) are used, due to the extended time needed for organoid establishment, growth, and maturation.^{172,173} Success rates also vary widely: for example, only $\sim 31\%$ of esophageal adenocarcinoma biopsies successfully yielded organoid lines,¹⁷⁴ and a recent meta-analysis in gastric cancer reported an average success rate of $\sim 66.6\%$ across multiple studies.¹⁷⁵ Such variability reflects both biological heterogeneity and technical attrition. They also indicate that only specific subtypes of the above-mentioned cancer entities may preferentially support success *in vitro* cultures, thereby introducing a bias in representability.

In practice, organoids remain valuable for cell banking or expansion when sample material is limited. Still, for truly time-sensitive applications, the most direct strategy is to introduce freshly collected patient tissue into a ToC system, thus bypassing the lengthy organoid-generation step. Even so, this approach faces significant challenges, including the technical complexity of microfluidic handling, the need for skilled personnel, and rigorous quality-control standards, factors that currently restrict clinical use to well-equipped research centers. Finally, the regulatory frameworks for validating and approving physical transport-based biomarkers remain far less mature than those for molecular diagnostics, posing a critical bottleneck to complete clinical translation.

4. Mechanisms of resistance: beyond cellular autonomy

Building on the CAF-driven causality framework established in section 2, we now examine how the spatial and temporal organization of the TME governs transport and contributes to therapy resistance beyond cell-autonomous mechanisms.

4.1 Temporal dynamics of transport and therapeutic windows

Time is a critical yet often underappreciated dimension of tumor biology. Microfluidic and theoretical studies consistently report biphasic transport behavior, characterized by an early advection-facilitated phase followed by diffusion-limited penetration as stromal remodeling progresses. Elliott and Yuan first formalized this transition, describing an initial permeability-dominated regime followed by transport restriction.¹⁷⁶ Similar phase-dependent dynamics have since



been reported for small molecules,¹⁷⁷ and immune-cell trafficking.¹⁷⁸ These observations challenge static views of drug delivery and highlight the need to consider temporal evolution when interpreting therapeutic efficacy.

During early tumor development, a transient degradative phase, often dominated by MMP activity, can increase matrix permeability and reduce transport resistance.¹⁷⁹ Drug delivery during this short window may result in relatively homogeneous penetration, even in confined tumor volumes. However, this permissive state is short-lived. As tumors mature, collagen deposition and enzymatic crosslinking progressively transform the ECM into a dense, transport-resistant network.³³ Simultaneously, vascular permeability declines and size-exclusion thresholds emerge, hindering diffusion even for small-molecules.^{3,180} Drug regimens that are effective during early stages may thus fail once stromal barriers are established.

Hypoxia plays a central role in orchestrating these transitions.¹⁸¹ Oxygen depletion activates hypoxia-inducible factor 1- α (HIF-1 α) signaling, upregulating MMPs while concurrently inducing ECM-modifying enzymes such as prolyl 4-hydroxylase α -subunit (P4HA), procollagen-lysine 2-oxyglutarate 5-dioxygenase (PLOD), and the LOX/LOXL family.^{31,181} Although hypoxia-driven angiogenesis transiently restores oxygenation, this relief ultimately favors matrix stabilization, increased IFP, and tissue stiffening.^{127,182} As oxygenation improves, metabolic shifts suppress acidic protease activity, further stabilizing the ECM.¹⁸³

These dynamics have been directly observed in tumor-on-chip systems. In PDAC-on-chip models, collagen accumulation and declining intrinsic permeability occurred primarily between days 7 and 21, and correlated with reduced gemcitabine efficacy.¹³⁴ Such observations illustrate that therapeutic windows are governed not only by drug properties but also by the kinetics of stromal remodeling, which vary across tumor types and patients.

4.2 Mechanical stress as a spatial regulator of transport

Tumor growth inherently generates mechanical stress through the combined effects of cellular proliferation, ECM deposition, and interstitial fluid accumulation within confined tissue. Biomechanically, solid tumors behave as biphasic materials, comprising a solid matrix and an interstitial fluid phase, whose coupled behavior is well described by poroelastic or poroviscoelastic models.

Collective CAF-mediated matrix contraction generates macroscopic solid stress in the kilopascal range (\sim 1.3–13 kPa) in human tumors.^{164,184} Matrix densification simultaneously reduces intrinsic permeability, and elevates IFP, leading to vessel compression and pore constriction across multiple length scales.^{29,34,164,180,185–187} These stresses are spatially heterogeneous and peak at the tumor-stroma interface, where transport resistance is maximal.^{164,186} As a result, drug and nutrient delivery becomes highly uneven, generating intratumoral transport heterogeneity.

4.3 Mechanotransduction and feedback amplification of resistance

Mechanical stress is not a passive consequence of tumor growth but an active regulatory signal. Through mechanotransduction pathways, stromal and tumor cells convert physical forces into transcriptional programs that reinforce resistance.^{30,44,188}

Tensile strain promotes integrin clustering and focal adhesion formation, activating focal adhesion kinase (FAK)-mediated signaling cascades that promote AKT-driven survival under stress.^{30,127,182} Matrix stiffening induces YAP/TAZ nuclear translocation, driving expression of ECM genes such as COL1A1/COL1A2, FN1, LOX, and connective tissue growth factor (CTGF).^{30,127,182,189,190} This establishes a positive feedback loop in which stiffening promotes further matrix deposition and crosslinking.

Importantly, mechanical cues also influence CAF plasticity. Chronic stress can induce transition between CAF phenotypes,^{9,55,191} thus complicating therapeutic strategies aimed at selectively targeting specific fibroblast subpopulations. These feedback mechanisms integrate mechanical heterogeneity, transport limitation, and stromal diversity into a self-sustaining resistance program.

4.4 Extrinsic resistance revealed by patient-derived transport phenotypes

Therapy resistance in solid tumors arises from both intrinsic (cell-autonomous) and extrinsic (transport-limited) mechanisms. While intrinsic resistance, such as drug-target mutations or enhanced drug efflux, has been extensively reviewed,⁴ extrinsic resistance arises from the physical constraints of the TME. Dense ECM deposition, enzymatic crosslinking, and elevated IFP can prevent drugs from reaching otherwise sensitive tumor cells.^{22,31,124,180,192}

Patient-derived tumor-on-chip systems provide direct experimental evidence for transport-limited resistance. By preserving native CAF composition and ECM architecture, these platforms reproduce clinically relevant barriers to drug penetration.¹⁹³ Across multiple tumor entities, effective permeability and matrix organization consistently emerge as dominant determinants of therapeutic response.^{124,194,195} Tumors with higher permeability respond faster and more strongly to treatment, whereas dense, CAF-rich matrices delay drug accumulation and blunt efficacy.

Several studies demonstrate the translational relevance of this transport phenotype. Multiplexed microfluidic platforms that culture intact tumor slices preserve native ECM architecture and stromal composition and generate spatially resolved drug-response readouts within days, directly linking tissue penetration to efficacy.¹⁹⁶ Patient-derived pancreatic tumor-on-chip systems incorporating organoids with stellate cells and macrophages recapitulate desmoplastic matrix deposition and show altered chemotherapy responses when stromal composition is experimentally modified, supporting a causal role for matrix-mediated transport barriers.¹⁹⁷ Fully



3D-printed and microfluidic tumor-on-chip devices sustaining patient spheroids or explants further demonstrate concordance with matched *in vivo* models, indicating translational relevance of transport-aware functional testing.¹⁹⁸ Complementary *in vivo* work in pancreatic cancer shows that enzymatic hyaluronan depletion reduces interstitial fluid pressure, restores vascular perfusion and increases intratumoral drug accumulation, converting a poorly perfused, drug-refractory tumor into a treatment-responsive one.^{124,199,200}

Together, these findings demonstrate that extrinsic resistance is not a fixed property but a modifiable consequence of transport limitation. ToC platforms therefore provide a functional complement to molecular biomarkers by revealing whether therapeutic agents can physically access their targets within the tumor microenvironment.

4.5 Scope and outlook

Despite these advances, clinical translation of transport-based phenotyping remains constrained by assay duration, technical complexity, and regulatory uncertainty. While organoids remain valuable for cell expansion, direct use of freshly resected tissue in ToC systems offers a more time-efficient route for assessing transport phenotypes, albeit with logistical challenges. Regulatory frameworks for physical, transport-based biomarkers are still less mature than those for molecular diagnostics, representing a key bottleneck for broader adoption.

Nonetheless, the convergence of mechanistic modeling, microfluidic experimentation, and patient-derived systems provides a strong foundation for targeting stromal barriers as a therapeutic strategy. The following section examines how these insights are being translated into interventions that actively modulate CAF-driven matrix remodeling to improve drug delivery.

Extended discussion of clinical validation studies, regulatory considerations, and practical constraints is provided in sections S9–S13.

5. Therapeutic strategies: targeting the physical microenvironment

5.1 Matrix normalization and mechanical modulation approaches

Therapies targeting the tumor stroma rather than the tumor cells usually aim to restore transport-permissive properties while preserving the tissue architecture. Because excessive matrix ablation can paradoxically accelerate tumor progression by releasing mechanical growth constraints, most current strategies focus on matrix normalization or selective modulation rather than complete removal.^{13,201}

In this context, stromal targeting is best viewed to rebalance transport rather than to eliminate the extracellular matrix.

Beyond enzymatic degradation, several pharmacological agents have demonstrated unexpected anti-fibrotic effects. Losartan, an angiotensin II receptor blocker, suppresses the expression of profibrotic mediators such as insulin-like growth factor-1 (IGF1), TGF β , CCL11 (eotaxin), MMP9, COL1A1, and thrombospondin (THBS2), thereby inhibiting pathological TGF- β -driven fibroblast activation.¹²⁵ By reducing collagen synthesis and crosslinking while sparing normal stromal homeostasis, Losartan exemplifies a normalization strategy that selectively targets tumor-associated fibrosis. Other approaches modulate tissue mechanics without disrupting ECM architecture. Halofuginone, initially developed for autoimmune diseases, interferes with multiple matrix-related signaling pathways, including TGF- β /Smad3/Nrf2, Akt/mTORC1/Wnt/ β -catenin, and HDAC2 signaling, reducing pathological matrix crosslinking and softening the stroma while preserving its structural integrity.²⁰² Similarly, rho-associated protein kinase (ROCK) inhibitors reduce actomyosin contractility, relieving vessel compression, improving, and enhancing drug delivery.^{203,204} Together, these approaches demonstrate that redistributing mechanical stress can improve transport as effectively as bulk matrix degradation.

Despite these benefits, stromal modulation carries inherent risks. Excessive increases in vascular permeability may inadvertently facilitate tumor cell dissemination or compromise local containment.²⁰⁵ Consequently, patient stratification based on stromal features such as HA content or tissue stiffness, together with careful risk–benefit assessment, is essential for safe clinical translation.¹²⁶

Additional stromal-modulation strategies are discussed in section S14.

5.2 Sequential therapy design

Sequential therapy strategies exploit the temporal dimension of stromal remodeling by conditioning the TME prior to cytotoxic or immunomodulatory treatment. Rather than co-administering all agents simultaneously, a first phase aims to reduce stromal resistance and improve transport, followed by a second phase that delivers the therapeutic payload under more permissive conditions. Preclinical studies have demonstrated that such conditioning significantly enhances intratumoral drug penetration and efficacy. For example, Zhang *et al.* (2024) designed a dendritic polymer system that initially delivered dasatinib-loaded nanoparticles to reprogram CAF metabolism and reduce collagen synthesis, followed by chemotherapeutic administration that achieved deeper penetration and greater tumor cell kill in the remodeled TME.³²

Sequential strategies are particularly compelling in immunotherapy. Reducing matrix stiffness or relieving physical barriers can restore CD8⁺ T-cells infiltration into tumor cores, effectively converting immune-excluded (cold) tumors into inflamed, therapy-responsive states. Preclinical studies have shown that stromal normalization enhances the



efficacy of immune checkpoint blockade by facilitating T-cell access to previously-excluded regions.^{206,207} Inhibition of LOX/LOXL-mediated collagen crosslinking follows the same principle, improving CD8⁺ T-cell infiltration and sensitizing tumors to anti-PD-L1 therapy.²⁰⁸

These approaches highlight the importance of timing and context: the therapeutic window for the second, effector phase is determined not only by drug pharmacokinetics but also by the dynamics of stromal remodeling. Defining this window requires imaging-guided assessment of transport and mechanical properties, reinforcing the relevance of transport phenotyping in therapy design.

Expanded examples of sequential and conditioning therapies are provided in section S15.

5.3 Clinical trial landscape and physical biomarkers

Several stromal-targeting strategies have progressed to clinical trials, revealing both their therapeutic potential and their limitations (Table S4). In PDAC, PEGPH20 trials showed that patient stratification by HA content is essential: clinical benefit was restricted to HA-high tumors, underscoring the need for pre-treatment biomarker selection.^{126,209}

Functional imaging is increasingly central to identifying patients likely to benefit from transport-targeted therapies and to monitoring treatment response. Magnetic resonance elastography (MRE) provides a non-invasive measure of tumor stiffness and has been shown to predict or stratify neoadjuvant response in breast cancer cohorts.²¹⁰ Dynamic contrast-enhanced MRI (DCE-MRI) offers complementary information on vascular function. The volume transfer constant (K^{trans}) integrates blood flow, vascular permeability, and surface area, with reduced K^{trans} values reflecting perfusion impairment caused by vessel compression or limited conductance. In PDAC, PEGPH20 treatment rapidly increased K^{trans} , correlating with HA depletion, vascular decompression, and improved drug delivery.^{211–213}

When applied longitudinally, these imaging modalities enable stratification of patients and response-adaptive therapy by aligning treatment decisions with improvements in perfusion or reductions in stiffness. However, broader clinical adoption remains limited by variability in imaging infrastructure, the absence of standardized physical biomarker thresholds, and safety concerns associated with stromal modulation. Extended clinical imaging and biomarker studies are summarized in section S16.

5.4 Integrating physical microenvironment targeting into precision oncology

Taken together, the strategies outlined above form a therapeutic continuum, ranging from matrix normalization and mechanical modulation to sequential conditioning protocols and imaging-guided adaptive therapies. Enzymatic degradation can induce rapid increases in permeability but requires stringent patient selection; mechanotherapeutics approaches achieve more gradual and reversible

normalization; and sequential strategies exploit transient windows of enhanced accessibility to boost the efficacy of cytotoxic and immunotherapies.

Successful clinical translation will depend on integrating robust physical biomarkers such as stiffness and perfusion into trial design and therapeutic decision-making. In this framework, physical microenvironment-targeted therapies complement molecular precision medicine by restoring drug accessibility and overcoming transport-limited resistance rather than by directly targeting tumor-cell-intrinsic vulnerabilities.

6. Critical assessment and future directions

6.1 Current consensus and open controversies

There is now broad agreement that fibroblast-driven ECM remodeling creates measurable transport limitations in desmoplastic tumors. Mechanistic, pathological, and biophysical studies converge on the view that increased matrix density and crosslinking elevate tissue stiffness while reducing intrinsic permeability, thereby limiting molecular and cellular access to malignant cells.^{3,31,182} These effects are particularly prominent in stroma-rich malignancies such as pancreatic ductal adenocarcinoma and selected breast and head-and-neck cancers, where the stromal compartment dominates tissue volume and mechanical behavior.

Importantly, the transport barrier is neither homogeneous nor static. Instead, tumors exhibit a mosaic of regions ranging from relatively compliant to highly compacted and crosslinked matrices, whose microscale organization governs local transport. ECM composition, alignment, porosity, and crosslinking dynamically evolve in response to CAF activity, therapy, and perfusion. These parameters are experimentally accessible, and can be measured, modeled, and even perturbed using microfluidic ToC platforms and complementary *in vivo* approaches.^{192,214} Representative ongoing clinical trials that explicitly or implicitly target physical and stromal transport barriers are summarized in Table S4.

Despite this consensus, several mechanistic controversies remain with direct translational implications. First, interstitial flow can exert opposing effects depending on its magnitude, geometry, and the size of the carried agents. Modest interstitial flow may enhance the advective delivery of macromolecules, whereas elevated flow can reverse transcapillary gradients and impair the extravasation of most therapeutics.^{215,216} This duality explains why identical interventions may improve the delivery of one agent while reducing the efficacy of another. Second, stromal targeting strategies, including CAF ablation, reprogramming, enzymatic degradation, or mechanical modulation, cannot be applied uniformly. Their efficacy depends on CAF subtype composition, spatial organization, ECM composition, and disease stage.²¹⁷ Consequently, stromal interventions must be matched to the tumor's biophysical phenotype, reinforcing



the need for individualized, transport-informed strategies rather than one-size-fits-all approaches. These debates are not merely conceptual: they directly shape trial design, patient stratification, and therapeutic sequencing.

6.2 Methodological, translational, and reporting gaps

Progress in quantitatively linking matrix remodeling to transport physics represents a conceptual shift in understanding therapeutic resistance. It extends intervention strategies beyond molecular targets to the physical infrastructure of tumors, recognizing stromal-mediated transport barriers as resistance mechanisms equivalent in importance to intrinsic cellular ones.

The causality framework linking fibroblast activation, ECM remodeling, and transport barrier formation provides mechanistic clarity for the design of rational interventions. By transforming descriptive observations into predictive

models, this framework identifies optimal intervention points within pathological cascades and supports evidence-based sequencing and combination strategies targeting both physical and molecular resistance mechanisms. Microfluidic platforms have evolved from proof-of-concept tools into validated experimental systems with increasing clinical relevance. By recapitulating patient-specific tumor architectures under controlled conditions, they bridge the gap between reductionist *in vitro* assays and complex *in vivo* systems. This progress is illustrated by the growing integration of commercial ToC technologies, such as Emulate's Zoë-CM2® culture module,^{218,219} InSphero's Akura™ organ-on-chip,^{220,221} and MIMETAS's OrganoPlate®,²²² into oncology pipelines, where they reproduce clinically relevant drug-response phenotypes more faithfully than conventional 2D assays.

Nevertheless, major gaps remain. Standardization of experimental protocols, reporting of transport parameters,

Clinical Translation Roadmap for Transport-Informed Precision Oncology (2024–2034)

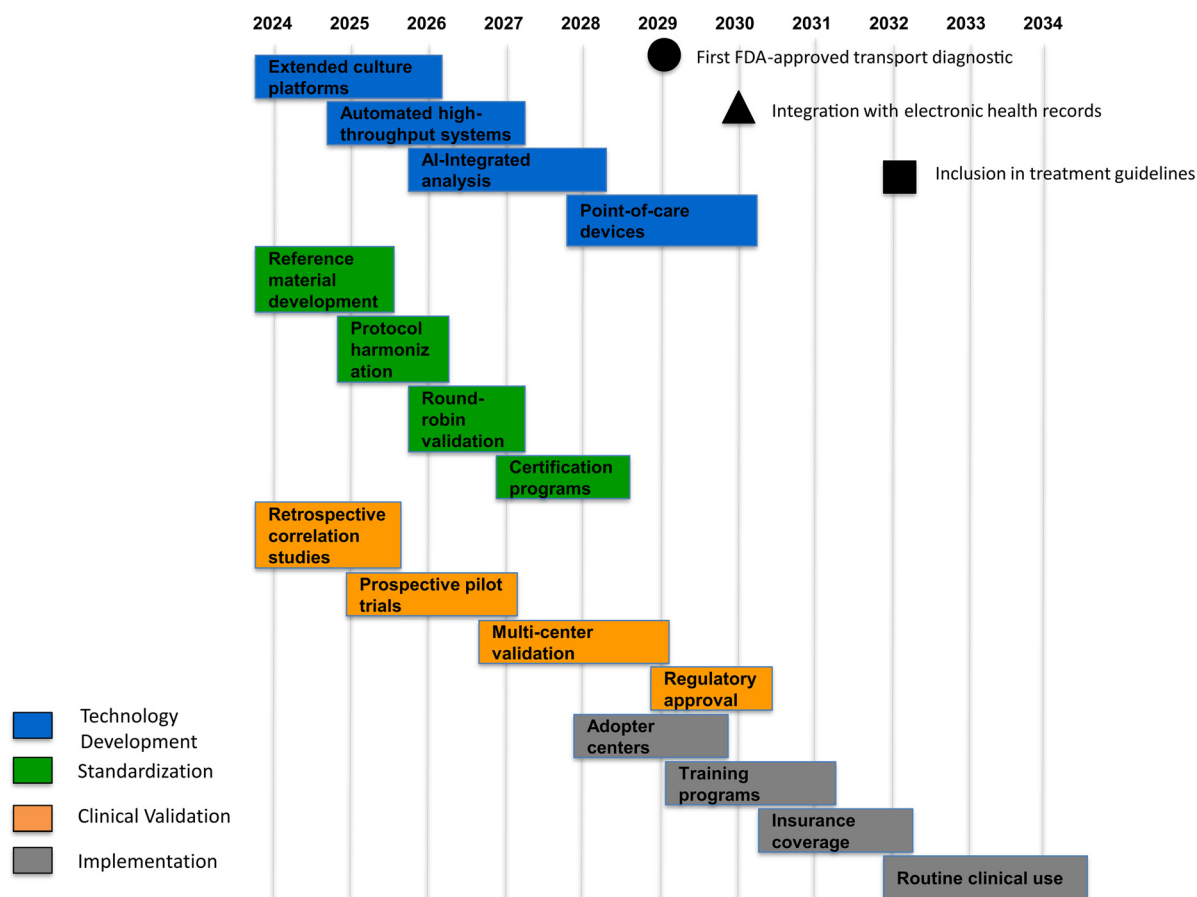


Fig. 3 Clinical translation roadmap for transport-informed precision oncology (2024–2034). Conceptual Gantt chart outlining a prospective, expert-informed roadmap for the clinical translation of transport phenotyping over a 10 year horizon. Four coordinated development tracks are shown: technology development, standardization, clinical validation, and implementation. Milestones are indicated by shape-coded markers: circle corresponds to the anticipated emergence of the first FDA-approved transport diagnostic (2029), triangle indicate expected integration with electronic health records (2030), and square represent potential inclusion in treatment guidelines (2032). Timelines are indicative and scenario-based, reflecting current technological maturity, regulatory trajectories, and published clinical precedents, and may vary depending on cancer type, regulatory context, and adoption pathways.



and definition of clinically actionable contexts of use are still lacking. A step-by-step overview of practical workflow constraints, key experimental decision points, and transport-relevant quality controls in patient-derived tumor-on-chip studies is provided in Table S5. Biological studies must increasingly treat physical properties as mechanistically equivalent to molecular determinants of tumor behavior, while regulatory frameworks must evolve to accommodate physical biomarkers under validation standards comparable to genomic markers.

6.3 Outlook: from transport barriers to therapeutic vulnerabilities

Within the next decade, transport profiling could complement genomic characterization in clinical decision-making. Patient stratification would then integrate both molecular alterations and drug-accessibility profiles, enabling sequential strategies that normalize physical properties before targeted delivery. The translation of transport phenotyping from experimental platforms to routine clinical use will require coordinated progress across technology development, standardization, clinical validation, and implementation. A prospective, scenario-based roadmap outlining these interdependent milestones over the next decade is summarized in Fig. 3. Mechanical modulation, stromal reprogramming, and externally applied physical interventions may become standard components of multimodal therapy. In this context, computational and AI-assisted approaches may further support transport phenotyping by guiding experimental design, parameter exploration, and sequential intervention strategies. Representative examples of such optional frameworks and their minimal data requirements are summarized in Table S6.

Beyond improving existing treatments, early intervention during matrix remodeling could prevent barrier formation

altogether. Detection of pre-barrier states may enable prophylactic normalization strategies, eliminating malignancies before transport limitation arises. These principles extend beyond oncology to other matrix-mediated pathologies such as fibrosis, atherosclerosis, and neurodegeneration. For patients with transport-limited malignancies refractory to current therapies, integrating transport physics with precision oncology offers a promising therapeutic avenue. Physical barriers that currently protect tumors may ultimately become exploitable therapeutic vulnerabilities.

7. Concluding remarks

This review highlights mass-transport phenotyping as a unifying framework for understanding how stromal architecture, mechanical confinement, and interstitial fluid dynamics jointly regulate therapeutic accessibility in solid tumors. Rather than treating biochemical, mechanical, and transport parameters as independent features, tumor-on-chip platforms enable these dimensions to be made experimentally commensurate and quantitatively comparable, thereby complementing molecular and genomic profiling with a functional, biophysical perspective on drug accessibility. By articulating the conceptual links between stromal remodeling and transport barriers, and by outlining practical priorities for platform design and reporting, we aim to support the development of tumor-on-chip systems that are both biologically meaningful and experimentally interoperable. To translate these principles into actionable design criteria, Table 1 summarizes practical development directions for next-generation ToC platforms explicitly dedicated to mass-transport phenotyping.

Such platforms are not intended to replace existing experimental models, but rather to serve as a complementary analytical layer. By enabling mechanistic questions related to drug, nutrient, and immune-cell access to be addressed

Table 1 Practical development directions for next-generation ToC platforms for mass-transport phenotyping

Capability	Why it matters?	Metrics/readouts
Controlled modulation of mechanical confinement	Tests how solid stress affects drug/oxygen/immune access	Applied compressive stress (kPa), strain, ECM porosity, cell viability maps
Independent tuning of interstitial pressure and perfusion	Distinguishes advective vs. diffusive transport contributions	ΔP (Pa), perfusion flow rate ($\mu\text{L min}^{-1}$), tracer velocity, perfusion fraction
ECM that supports physiologically relevant remodeling	Recapitulates desmoplasia and dynamic barrier formation	Time-course of ECM density, fiber alignment (SHG), crosslink markers, stiffness (kPa)
Non-invasive readouts of transport and mechanics	Enables longitudinal experiments without perturbation	FRAP, fluorescent tracer diffusion (D_{eff}), microbead-based or elastography stiffness maps, pressure sensors
Device materials compatible with quantitative assays	Avoids artefactual adsorption or loss of analytes	Adsorption tests (percent recovery), surface treatment documented, control tracer mass balance
Compatibility with standard cell-culture workflows	Promotes adoption and reproducibility	Multi-well footprint, pipet access, common plate reader/microscope compatibility
Transport-centered reporting standards	Enables cross-platform, quantitative comparison	Minimum dataset: ECM stiffness, k , D_{eff} , ΔP , tracer properties, device geometry, and access to the analysis code used to compute transport parameters



quantitatively, transport-aware ToC systems provide an intermediate scale between reductionist *in vitro* assays and complex *in vivo* models, where physical constraints often remain difficult to disentangle. Ultimately, adopting mass-transport-aware ToC approaches offers the opportunity to move beyond descriptive observations of treatment resistance toward a predictive, parameter-based understanding of tumor response.

This shift has the potential to refine experimental strategies, guide the rational design and sequencing of therapies, and improve the interpretability and reproducibility of preclinical studies across laboratories. For transport-limited malignancies that remain refractory to current treatments, integrating transport physics into precision oncology may help transform physical barriers from passive obstacles into actionable therapeutic vulnerabilities.

Glossary

Cancer-associated fibroblasts (CAFs): activated stromal fibroblasts within tumors that produce and remodel ECM. Heterogeneous population including matrix-producing (mpCAFs) and matrix-remodeling (mrCAFs) subtypes.

Intrinsic permeability (k): a geometric property of a porous medium that quantifies its ability to transmit fluid independently of the fluid viscosity. It is expressed in m^2 and depends on ECM porosity, pore size, fiber alignment, tortuosity, and matrix compaction. In tumors, k typically decreases by several orders of magnitude during CAF-driven desmoplastic remodeling, directly limiting interstitial flow and advective transport.

Fluid viscosity (μ): dynamic viscosity of the interstitial fluid (Pa s), assumed close to that of aqueous culture media and treated as constant in this review. Variations in advective transport are therefore attributed primarily to ECM-controlled changes in intrinsic permeability k .

Effective diffusivity (D_{eff}): apparent diffusion coefficient of a solute within a tissue or ECM, accounting for porosity, tortuosity, binding, and confinement. D_{eff} is lower than the free diffusivity in solution (D_0) and dynamically evolves with matrix remodeling and solid stress.

Interstitial fluid pressure (IFP): pressure within the fluid-filled spaces between cells and matrix fibers. Elevated in tumors (1–8 kPa) compared to normal tissues (–1–+1 kPa), reducing transcapillary pressure gradients and limiting advective transport despite intact vasculature.^{31,223}

Porosity (ϕ): fraction of tissue volume occupied by fluid-filled voids. In tumors, porosity decreases with ECM densification and compression, directly reducing diffusivity and intrinsic permeability.

Mechanotransduction: the process by which cells convert mechanical stimuli into biochemical signals. Key pathways involved include integrin clustering, focal adhesion kinase (FAK) activation, and the nuclear translocation of YAP/TAZ.

Solid stress: compressive and tensile forces generated within growing tumors as cells proliferate against

surrounding tissue constraints. It can reach 1.3–13.0 kPa, sufficient to collapse blood vessels.^{164,184}

Tortuosity: dimensionless parameter describing the geometric complexity of transport pathways in a porous medium. In biological tissues, higher tortuosity reflects elongated and convoluted diffusion or flow paths imposed by ECM architecture. It is commonly defined as $\lambda = \sqrt{D_0/D_{\text{eff}}}$, where D_0 is the free diffusivity and D_{eff} the effective diffusivity in tissue.

Mass-transport phenotyping: functional characterization of tumors based on quantitative transport properties (e.g., diffusivity, permeability, interstitial flow) that determine drug, nutrient, and immune-cell accessibility, independently of molecular target expression.

Tumor-on-chip (ToC): microfluidic *in vitro* platforms that recapitulate key aspects of tumor architecture, perfusion, and stromal remodeling under controlled conditions, enabling quantitative analysis of transport and mechanical phenomena.

Extrinsic (transport-limited) resistance: apparent therapeutic resistance arising from limited drug delivery due to physical barriers in the tumor microenvironment, despite intrinsic cellular sensitivity to the drug.

Author contributions

Doriane Le Manach: conceptualization, visualization, project administration, writing – original draft, writing – review & editing. Vincent Senez: conceptualization, funding acquisition, supervision, review & editing. Matthias Nees: funding acquisition, resources, supervision, review & editing.

Conflicts of interest

There are no conflicts to declare.

Data availability

No primary research results, experimental data, software or code have been included, and no new data were generated or analyzed as part of this review.

Supplementary information: Section S1 and Table S1 provide a comprehensive list of causal input parameters, transport-related outputs, symbols, units, and corresponding experimental readouts for CAF-driven transport phenotyping. (Sections 2.1 & 2.2); **Sections S2 and S3** detail experimental, computational, and interventional strategies for calibrating transport parameters and modulating CAF-driven matrix barriers within transport-phenotyping frameworks. (Sections 2.3 & 2.4); **Fig. S1** presents a hierarchical overview of tumor model systems used for mass-transport studies, positioning tumor-on-chip platforms relative to conventional *in vitro* and *in vivo* models. (Section 3.1); **Section S4 and Table S2** provide a comparative overview of microfluidic device structural materials, surface functionalization strategies, and their implications for quantitative transport assays. (Sections 3.1 &



3.2); **Sections S5 and S6**, together with **Table S3**, summarize hydrogel and ECM choices, matrix stabilization strategies, and confinement approaches for CAF-driven transport phenotyping, including common experimental pitfalls. (Sections 3.2 & 3.3); **Section S7** discusses measurement techniques, technical limitations, and multimodal strategies for quantifying diffusion, advection, and matrix remodeling in tumor-on-chip systems. (Section 3.4); **Section S8** outlines recommended minimum reporting standards and reproducibility considerations for transport-resolved tumor-on-chip studies. (Section 3.5); **Sections S9–S13** address practical, regulatory, and translational constraints for transport-based phenotyping, including representability, success rates, validation examples, and comparison with molecular biomarkers. (Section 4.5); **Sections S14–S16** expand on stromal-targeting strategies, sequential therapy concepts, and imaging biomarkers relevant to transport-guided therapeutic interventions. (Sections 5.1, 5.2 & 5.3); **Table S4** summarizes representative ongoing clinical trials targeting physical and stromal barriers to improve drug transport in solid tumors. (Section 6.1); **Tables S5 and S6** detail workflow constraints for patient-derived tumor-on-chip transport phenotyping and optional AI-assisted experimental design strategies (Sections 6.2 & 6.3). See DOI: <https://doi.org/10.1039/d5lc01131k>.

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