

# The Physics of Immune Cooperation: Dimensional Surveillance and Attractor Enforcement in Multicellular Systems

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## Abstract

The immune system is conventionally understood as a pattern-recognition system that distinguishes “self” from “non-self” via molecular signatures. We propose a more fundamental interpretation: immunity is **dimensional surveillance**—the detection and elimination of cells whose dynamical complexity has collapsed below tissue-appropriate thresholds. Cancer cells, virally-infected cells, and senescent cells share a common signature: reduced effective dimensionality, reflecting escape from the high-dimensional organismal attractor into simpler replicative or dysfunctional states. Immune receptor interactions (CD molecules, MHC-TCR coupling) function as synchronization probes that measure the dynamical complexity of target cells. Inflammation is controlled destabilization—a transient increase in tissue volatility that enables

state transitions (clearance, healing, remodeling). The HPA axis modulates detection thresholds: acute stress raises tolerance transiently (survival prioritization), while chronic stress produces persistently shallow attractor basins with elevated thresholds, enabling defector cells to escape surveillance. We formalize this using Price equation multilevel selection: within-organism selection (immune enforcement against low-D defectors) trades off against between-organism selection (survival under resource constraints). Finally, we demonstrate that immune surveillance entails a thermodynamic cost: “dynamical friction” from coupling to low-dimensional targets drives metabolic exhaustion, explaining why chronic antigen exposure leads to T-cell functional collapse. When the sensor itself collapses, distinct target states alias onto the same low-dimensional projection, producing either indiscriminate attack (autoimmunity) or indiscriminate tolerance (exhaustion)—explaining the paradox of inflamm-aging. This framework unifies cancer immunology, autoimmunity, chronic inflammation, and the evolution of adaptive immunity under a single dynamical principle: immunity is cooperation enforcement, operating through dimensional measurement at the cellular scale.

**Keywords:** immunity, dimensional surveillance, attractor dynamics, inflammation, cooperation, multilevel selection, cancer immunology, HPA axis

## 1 Introduction: What Is the Immune System Actually Measuring?

The textbook account of immunity centers on molecular recognition. Innate immunity detects conserved pathogen-associated molecular patterns (PAMPs); adaptive immunity generates diverse receptors that recognize specific antigens. The self/non-self distinction is implemented through negative selection in the thymus and tolerance mechanisms that prevent attack on host tissues.

This framework works well for infectious disease but struggles with cancer. Cancer cells are genetically “self”—they carry the host genome, express host proteins, and display host

MHC molecules. The field has responded by adding epicycles: “danger signals,” damage-associated molecular patterns (DAMPs), stress ligands, and tumor-associated antigens. Each explains some observations but none provides a unified mechanism.

We propose that the immune system is measuring something more fundamental than molecular identity: it is measuring **dynamical complexity**. Cells embedded in tissue perform complex, context-sensitive computations—responding to neighbors, integrating signals, maintaining tissue architecture. This complexity manifests as high effective dimensionality ( $D_{\text{eff}}$ ) in the cell’s dynamical state space. Indeed, Cohen et al. [11] have argued that aging itself is best understood as a progressive loss of structural and functional complexity, where systems collapse from high-dimensional homeostatic states into simpler, dysregulated attractors. Cells that have “defected” from the tissue program—whether through malignant transformation, viral hijacking, or senescent collapse—exhibit reduced dimensionality: they have fallen into simpler attractors.

Immune surveillance, on this view, is dimensional surveillance. The receptor-ligand interactions that immunologists study (CD4, CD8, MHC-TCR, checkpoint molecules) function as **synchronization probes**: the immune cell couples its oscillatory dynamics to the target cell and measures the response. High-dimensional targets produce complex, context-appropriate coupling; low-dimensional targets produce stereotyped, simplified responses.

This reframing has immediate consequences:

1. **Cancer immune evasion** is not primarily about hiding antigens—it is about faking high-dimensional signatures (e.g., checkpoint expression that mimics normal tissue coupling).
2. **Autoimmunity** represents miscalibrated thresholds: the system attacks normal cells that transiently dip in dimensionality (stress, infection, metabolic challenge).
3. **Chronic inflammation** is the combination of shallow attractor basins (cells prone to dimensional collapse) and raised detection thresholds (immune tolerance under persistence).

tent stress)—defectors escape because the system is both producing more of them and catching fewer.

4. **Immunotherapy** works when it restores dimensional surveillance: checkpoint blockade removes the “fake high-D” signals that tumors use to evade detection.

## 1.1 The Evolutionary Logic

Multicellularity creates a cooperation problem. Individual cells “benefit” (in the short term) from defecting to a replicative program—this is cancer. The organism requires mechanisms to detect and eliminate defectors. But detecting defection is hard: defectors carry the same genome and can express the same surface proteins as cooperators.

The solution is behavioral surveillance. Cooperating cells exhibit high-dimensional dynamics because tissue function *requires* complex, context-sensitive behavior. Defecting cells exhibit low-dimensional dynamics because replication is a simple program. The immune system evolved to detect this dimensional signature.

This is a multilevel selection problem, formalizable via the Price equation:

$$\Delta\bar{z} = \underbrace{\text{Cov}(W_i, z_i)}_{\text{within-organism selection}} + \underbrace{E[W_i\Delta z_i]}_{\text{transmission bias}} \quad (1)$$

where  $z$  is a trait (e.g., replicative rate),  $W_i$  is fitness, and the covariance term captures selection among cells within an organism. Immune enforcement acts on this term: by eliminating high-replication (low-D) cells, the immune system suppresses within-organism selection for defection.

The cost is metabolic: maintaining immune surveillance requires energy. The benefit is organismal coherence: suppressing defectors preserves the high-dimensional dynamics required for tissue function. The tradeoff is modulated by the HPA axis, which adjusts surveillance intensity based on organismal state.

## 2 Dimensional Collapse as Defection

### 2.1 What Dimensionality Means for Cells

A cell’s “effective dimensionality” ( $D_{\text{eff}}$ ) refers to the number of independent degrees of freedom in its dynamical state. A cell performing complex tissue functions—responding to gradients, communicating with neighbors, adjusting metabolism to context—occupies a high-dimensional attractor. Its state trajectory explores many dimensions of the available state space.

A cell that has collapsed into a replicative program occupies a low-dimensional attractor. The cell cycle is a limit cycle: a stereotyped sequence of states that repeats. Viral infection similarly reduces dimensionality: the cell’s dynamics are dominated by the viral replication program. Senescence is another low-dimensional state: the cell has exited the complex tissue attractor but not entered replication. As noted by Cohen et al. [11], senescence represents a discrete “attractor state”—a deep basin that traps cells once they cross a critical threshold of dysregulation. We extend this by proposing that immune surveillance is the mechanism responsible for detecting cells that have fallen into these low-complexity basins.

Formally, consider a cell’s state as a vector  $h(t) \in \mathbb{R}^m$  evolving according to:

$$\frac{dh}{dt} = f(h, e, c) \tag{2}$$

where  $e$  represents environmental/tissue signals and  $c$  represents cell-autonomous programs. The effective dimensionality is the number of significant principal components of the trajectory  $h(t)$  over a characteristic timescale.

For a cell embedded in functional tissue:

- $e$  is high-dimensional (complex tissue signals)
- The attractor is high-dimensional (many modes active)
- $D_{\text{eff}} \sim O(10^2)$  or higher

For a cancer cell:

- $e$  is ignored or overridden (cell-autonomous replication)
- The attractor is low-dimensional (cell cycle limit cycle)
- $D_{\text{eff}} \sim O(10^0 - 10^1)$

The immune system, we propose, measures this difference.

## 2.2 Operationalizing Effective Dimensionality

While  $D_{\text{eff}}$  is a theoretical construct, several empirical metrics can approximate it:

- **PCA Participation Ratio.** Given eigenvalues  $\lambda_i$  of the state covariance matrix:

$$PR = \frac{(\sum_i \lambda_i)^2}{\sum_i \lambda_i^2} \quad (3)$$

This equals 1 if variance is concentrated in one dimension; it equals  $n$  if variance is uniformly distributed across  $n$  dimensions.

- **Spectral Entropy.** Given power spectrum  $P(f)$  normalized to a probability distribution:

$$H_{\text{spectral}} = - \sum_f P(f) \log P(f) \quad (4)$$

Low entropy indicates few dominant frequencies (low-D); high entropy indicates distributed power (high-D).

- **Transcriptional Heterogeneity.** In single-cell RNA sequencing, cells with complex tissue functions show higher transcriptional entropy than cells locked in replicative programs.

These metrics are not  $D_{\text{eff}}$  itself—they are projections of the underlying dynamical complexity onto observable quantities. But they provide testable predictions: cancer cells should

show lower participation ratios, lower spectral entropy, and lower transcriptional heterogeneity than matched normal tissue.

A caveat: high dimensionality per se is not sufficient for health. Pathological states like seizures and cytokine storms involve high-dimensional dynamics that are *uncoordinated*—high variance but low functional integration. The healthy regime is high-dimensional *and* coherent: many active modes, but modes that are coupled to tissue-level function. We might call this “functional dimensionality” to distinguish it from mere noise. The immune system, we suggest, detects the absence of coordinated complexity, not just the absence of complexity.

### 2.3 Evidence for Dimensional Collapse in Cancer

The “hallmarks of cancer” [1] can be reinterpreted as signatures of dimensional collapse:

1. **Sustaining proliferative signaling:** The cell has locked into a replicative attractor and no longer requires external signals to maintain it.
2. **Evading growth suppressors:** The cell no longer responds to tissue-context signals that would modulate its state.
3. **Resisting cell death:** The cell has exited the attractor basin that includes apoptosis as a response to damage.
4. **Enabling replicative immortality:** The cell cycle limit cycle has become indefinitely stable.
5. **Inducing angiogenesis and activating invasion:** Secondary adaptations that support the low-D replicative state.
6. **Genome instability:** A consequence, not a cause—the low-D attractor tolerates genomic variation because the replicative program is robust.

Crucially, the Mintz and Illmensee experiments [2] demonstrated that this collapse is reversible: teratocarcinoma cells injected into mouse blastocysts contributed to normal tissues, including functional sperm. The cells were not genetically “broken”—they were in a different attractor. The embryonic environment pushed them back into the high-dimensional tissue attractor.

## 2.4 Viral Infection as Dimensional Hijacking

Viral infection produces a similar dimensional signature. The virus commandeers cellular machinery for replication, collapsing the cell’s dynamics into a viral-production program. The infected cell no longer performs tissue-appropriate computation; it performs viral computation.

Crucially, a virus is not merely a packet of molecular information—it is a **template for an attractor**. When a virus enters a cell, it does not simply inject instructions; it captures the cell’s dynamics into a viral attractor basin. The viral genome encodes not just proteins but a dynamical program that, once initiated, becomes self-sustaining. The infected cell has “fallen into” the viral attractor.

This explains why the immune system uses similar mechanisms for cancer and infection: both are detected via the same dimensional signature. The difference is that infected cells also present viral peptides on MHC, providing an additional detection channel. But the dimensional collapse is primary; antigen presentation is secondary.

## 2.5 Prions as Protein-Level Attractors

Prions extend this logic to the molecular scale. A prion is not an organism or even a genome—it is a **protein conformation that constitutes a stable attractor**. The misfolded prion protein ( $\text{PrP}^{\text{Sc}}$ ) is thermodynamically stable and, critically, it templates the misfolding of normal  $\text{PrP}^{\text{C}}$  proteins it contacts.

In dynamical terms: the prion conformation is a low-dimensional attractor in protein

folding space. It propagates by pulling other proteins into the same basin. This is “infection” without any genetic material—purely attractor dynamics at the molecular level.

The immune system struggles with prions precisely because there is no dimensional collapse at the cellular level to detect. The cells producing misfolded proteins may still exhibit normal high-D dynamics; only the proteins themselves have fallen into a pathological attractor. This represents a blind spot in dimensional surveillance: the signal operates at the wrong scale.

Notably, humans show evidence of evolutionary adaptation to prion exposure. The Fore people of Papua New Guinea, who practiced ritualistic cannibalism, suffered a devastating epidemic of kuru (a prion disease) that at its peak killed 35 per 1000 population annually [8]. This created intense selection on the PRNP gene: survivors were predominantly heterozygous at codon 129, and a novel protective variant (G127V) appeared that provides near-complete resistance to prion infection [8, 9]. Worldwide PRNP haplotype diversity suggests similar selection pressures occurred in other human populations—implying prehistoric prion exposure, possibly through cannibalism, was more widespread than often assumed [10].

From an attractor perspective, the evolutionary response to prions is instructive: because the immune system cannot detect the protein-level attractor, selection instead modified the protein’s folding landscape itself. The G127V variant changes the prion protein’s conformational dynamics, making it harder to fall into the pathological attractor basin. When surveillance fails, the alternative is to engineer the landscape.

## 3 Immune Receptors as Synchronization Probes

### 3.1 The Coupling Hypothesis

Immune cells interact with target cells through multiple receptor-ligand pairs: TCR-MHC, CD4/CD8 co-receptors, costimulatory molecules (CD28-B7), coinhibitory molecules (PD-1-PD-L1, CTLA-4-B7), adhesion molecules (LFA-1-ICAM), and many others. The textbook

interpretation is that these provide recognition signals: “this is foreign” or “this is self.”

We propose a different interpretation: these receptor interactions implement **dynamical coupling**. The immune cell synchronizes its intracellular oscillators (calcium waves, signaling cascades, cytoskeletal dynamics) with the target cell and measures the coupling response.

A high-dimensional target cell produces complex, context-sensitive coupling: the signaling response depends on the cell’s current state, recent history, and tissue context. The immune cell’s probes return rich, variable information.

A low-dimensional target cell produces stereotyped coupling: the replicative program dominates all responses, so the immune cell’s probes return simplified, repetitive information.

The immune cell integrates this information over time and makes a decision: attack or tolerate. The decision threshold is modulated by systemic state (inflammation, cortisol, checkpoint signals).

### 3.2 Checkpoint Molecules as Dimensional Camouflage

Checkpoint molecules (PD-L1, CTLA-4 ligands) are conventionally understood as “brakes” on immune activation—signals that say “don’t attack this cell.” Many tumors upregulate these molecules to evade immunity.

In our framework, checkpoint molecules function as **dimensional camouflage**: they inject complexity into the coupling response, making a low-D cell appear high-D to the probing immune cell. The tumor is not hiding its antigens; it is faking its dynamical signature.

This explains why checkpoint blockade works: removing PD-1/PD-L1 signaling strips away the camouflage, revealing the tumor’s true (low) dimensionality. The immune system can then recognize and attack.

It also explains why checkpoint blockade causes autoimmunity: by removing a camouflage mechanism, we lower the effective threshold for dimensional detection. Normal cells that transiently dip in dimensionality (stress, infection, metabolic challenge) are now attacked.

### 3.3 Dissolving the Combinatorial Explosion

A classic puzzle in immunology is how the immune system handles combinatorial explosion.

The textbook framing:

- The space of possible foreign antigens is vast ( $\sim 10^{15}$  possible peptides).
- The repertoire of T cell receptors is large but finite ( $\sim 10^7\text{--}10^8$  unique clonotypes).
- How does the system “cover” antigen space with so many fewer receptors?

Standard answers invoke V(D)J recombination generating diversity, cross-reactivity allowing one receptor to recognize multiple antigens, and negative selection removing self-reactive clones. But these answers are unsatisfying—they describe the system without explaining how it actually solves the recognition problem with such apparent combinatorial mismatch.

Dimensional surveillance dissolves this puzzle. The immune system is not pattern-matching in a discrete antigen space; it is measuring a continuous property (dynamical complexity) of target cells. Consider:

1. **You don't need to match every antigen.** If the signal is “this cell has low-dimensional dynamics,” then any low-D cell triggers the response, regardless of its specific molecular makeup. There is no need to have a receptor for every possible antigen.
2. **Cross-reactivity is expected, not puzzling.** A receptor that measures dynamical coupling will respond similarly to many different low-D cells, because low-dimensionality itself is the conserved signal. “Cross-reactivity” is simply what dimensional detection looks like from a molecular perspective.
3. **Receptor diversity measures different aspects of complexity.** The vast diversity of TCRs is not about covering antigen space—it is about having diverse synchronization

probes that sample different dimensions of the target cell’s dynamics. Each receptor-MHC interaction tests a different aspect of the cell’s response; the ensemble provides a high-dimensional probe.

4. **Negative selection removes high-affinity self-responders.** In our framework, negative selection eliminates receptors that strongly couple to normal high-D self dynamics. What remains are receptors calibrated to detect the *absence* of normal complexity—exactly what dimensional surveillance requires.

The combinatorial explosion was never a real problem; it was an artifact of thinking about immunity as pattern-matching in sequence space rather than dimensional measurement in dynamical space. A thermometer does not need to “recognize” every possible temperature—it measures a scalar property. The immune system, we propose, detects anomalies in dynamical complexity—not by “measuring” in any cognitive sense, but through the mathematics of high-dimensional interaction.

### 3.4 Thymic Selection as Dimensional Filtering

The thymus is where T cell receptors are filtered before release into the periphery. The standard account: positive selection ensures TCRs can bind MHC, negative selection eliminates TCRs that bind self-peptides too strongly. But this framing is puzzling—how does eliminating self-reactive cells prepare the repertoire to recognize unknown foreign antigens?

Dimensional surveillance reframes thymic selection:

- **Positive selection** ensures the TCR can *couple dynamically* to normal tissue cells. A receptor that cannot synchronize with high-D self dynamics is useless as a probe—it would trigger on everything or nothing. Positive selection calibrates coupling strength.
- **Negative selection** eliminates TCRs that couple *too strongly* to high-D self. These would trigger chronic activation in normal tissue. What survives are receptors that couple moderately to high-D self but strongly to low-D deviations.

- **The final repertoire** is a collection of probes optimized to detect *absence* of normal complexity. Each TCR tests a different aspect of the target cell’s dynamics; the ensemble provides comprehensive coverage.

This explains why self-tolerance is not “learning” a list of self-antigens. The thymus trains receptors to recognize a *dynamical signature*—the high-dimensional, tissue-coupled complexity of normal cells. Anything that deviates from this signature triggers suspicion, regardless of its molecular identity.

### 3.5 High Dimensionality Is Not a Metaphor

There is a persistent false dichotomy in biology: either a system is “just random molecules” (acceptable) or it is “intelligent” (anthropomorphizing, unacceptable). High-dimensional dynamics offers a third option that dissolves this dichotomy.

Consider V(D)J recombination. The textbook description: random recombination of Variable, Diversity, and Joining gene segments generates antibody diversity. The numbers are astronomical—roughly  $10^{11}$  possible heavy chain combinations, multiplied again by light chain diversity. This is typically framed as “random” in the sense of noise, undirected, purposeless.

But high-dimensional randomness is qualitatively different from low-dimensional randomness. When a system samples randomly from  $10^{11}$  possibilities, it is not producing noise—it is *exploring a high-dimensional space*. The mathematical properties of high-D random sampling are precisely what enable function:

- **Coverage.** Random sampling in high-D space covers more of the space than intuition suggests. The probability that a random antibody will bind *something* with moderate affinity is high, because high-D spaces have more “surface area” relative to volume.
- **Projection.** Any low-dimensional structure embedded in high-D space will be intersected by random high-D sampling. A pathogen, being a low-D replicator, cannot hide

from a high-D search.

- **Refinement.** Somatic hypermutation and clonal selection then perform gradient descent in this high-D space, finding high-affinity variants. But crucially, the initial random coverage is what makes this refinement possible.

This is not a metaphor. The immune system does not “act as if” it were exploring high-dimensional space—it *is* exploring high-dimensional space. V(D)J recombination literally generates a high-dimensional distribution over sequence space. Clonal selection literally performs optimization in this space. The “intelligence” of the immune system—its ability to recognize novel pathogens, to remember past infections, to distinguish self from non-self—emerges from these high-dimensional dynamics.

The reductionist instinct is to say: “But it’s just molecules binding to molecules.” Yes—but the dynamics of those molecules embody computation in a deeper sense than discrete arithmetic. The immune system is not performing matrix operations; it is a continuous physical process with uncountably many degrees of freedom. Any finite-dimensional mathematical description is a projection, a map of a territory that contains more information than any map can capture. When we say the immune system “detects” low-dimensional dynamics, we mean that high-dimensional physical systems naturally couple differently to low-dimensional versus high-dimensional targets. This is physics, not arithmetic—the dynamics *are* the computation, not a representation of computation.

The philosopher Sidney Morgenbesser, responding to B.F. Skinner’s behaviorist elimination of mental vocabulary, reportedly asked: “Let me see if I understand your thesis. You think we shouldn’t anthropomorphize people?” The same retort applies here. If high-dimensional dynamics is what produces cognition in humans, then recognizing high-dimensional dynamics in immune systems is not anthropomorphizing—it is recognizing the same mathematics at work. We *are* molecular systems. The question is not whether molecules can compute, but what kinds of computation different molecular architectures perform.

This has implications beyond immunology. Wherever we see biological systems generating vast combinatorial diversity—neural connectivity, gene regulatory networks, microbial ecosystems—we should ask whether the high-dimensionality itself is functional, not merely noise to be averaged over. The dimension *is* the mechanism.

### 3.6 Computational Demonstration

To test whether dimensional complexity is actually discriminable, we simulated cellular dynamics as coupled oscillators with varying numbers of active modes. Low-dimensional “pathogen-like” cells were modeled with 2–5 oscillatory modes; high-dimensional “self-like” cells with 20–50 modes. We then asked: can an observer distinguish these populations based on temporal dynamics alone?

The answer is yes, decisively. Spectral entropy—a measure of how many frequency components contribute to a signal—discriminates low-D from high-D dynamics with Cohen’s  $d = 3.70$  and classification AUC = 0.998 (Figure 1). Low-dimensional systems produce coherent, periodic signals dominated by a few frequencies; high-dimensional systems produce complex, aperiodic signals with power distributed across many frequencies.

Crucially, this discrimination requires *temporal* information. Static snapshots—projections onto fixed basis vectors—do not reliably distinguish dimensionality because normalization equalizes total variance. The information lives in the time domain: how the system evolves, which frequencies it uses, how quickly correlations decay. This suggests that immune surveillance, if it operates on dimensional principles, must involve dynamic probing—sustained interaction between receptor and target, not instantaneous binding events.

The simulation also confirms the costly signalling prediction: complexity and replication trade off (Figure 2). Cells that invest resources in maintaining high-dimensional dynamics have less available for replication ( $r = -0.59$ ,  $p < 0.001$ ). This tradeoff ensures that dimensional complexity is an honest signal—it cannot be faked without paying the metabolic cost.

## 4 Costly Signalling: Why Dimensional Complexity Cannot Be Faked

### 4.1 The Honest Signal Problem

A critical question arises: if dimensional complexity is the signal, why can't defectors simply fake it? Pathogens routinely evolve molecular mimicry—expressing host-like surface proteins to evade recognition. Why can't cancer cells evolve to produce high-dimensional dynamics while maintaining their replicative advantage?

The answer lies in costly signalling theory [6, 7]. A signal is honest when faking it costs as much as producing it genuinely. The peacock's tail works as an honest signal of fitness because growing a large tail requires resources that unhealthy individuals cannot spare. There is no cheap way to fake it.

**Dynamical complexity is a costly signal.** Producing high-dimensional, context-sensitive dynamics requires:

1. **Metabolic investment:** Maintaining complex intracellular signaling networks, ion gradients, and regulatory cascades consumes ATP.
2. **Tissue integration:** Responding appropriately to neighbors requires actually sensing and processing their signals—not possible for a cell focused on autonomous replication.
3. **Coordination overhead:** High-D dynamics emerge from coupling to the tissue program. A cell running its own replicative program cannot simultaneously run the tissue program.
4. **Replication opportunity cost:** Resources invested in complexity are resources not invested in division. There is a fundamental tradeoff.

## 4.2 The Replication-Complexity Tradeoff

We can formalize this as a resource allocation problem. Let a cell have total resource budget  $R$ . It allocates fraction  $\alpha$  to complexity maintenance (signaling networks, tissue integration) and  $(1 - \alpha)$  to replication machinery. Then:

$$D_{\text{eff}} \approx f(\alpha) \quad (\text{increasing in } \alpha) \quad (5)$$

$$\text{Replication rate} \approx g(1 - \alpha) \quad (\text{increasing in } 1 - \alpha) \quad (6)$$

A cell cannot maximize both. The tradeoff is fundamental.

This tradeoff has deep evolutionary roots. High-dimensional dynamics require sustained energy expenditure—maintaining ion gradients, protein turnover, active transport, coordinated signaling cascades. Aerobic respiration produces roughly 15 times more ATP per glucose than anaerobic glycolysis. It is not coincidental that complex multicellular life appears in the fossil record only after the Great Oxidation Event, and that the Cambrian explosion of animal body plans follows the rise of atmospheric oxygen to near-modern levels. Oxygen didn't just enable larger bodies; it enabled the metabolic rates necessary for complex, high-dimensional cellular dynamics. The immune systems that enforce cooperation could only evolve once the energy budget existed to make dimensional complexity affordable—and to make its absence detectable.

We simulated this tradeoff using a coupled oscillator model where cells allocate resources between oscillator coupling (complexity) and replication rate (Figure 2). The results confirm strong negative correlation ( $r \approx -0.6$ ) between probed dimensionality and replication rate. Cells attempting “cheap camouflage” (adding noise without true investment) are detected; cells achieving “costly camouflage” (actually investing in complexity) lose their replicative advantage.

### 4.3 Why Checkpoint Camouflage Works (Temporarily)

Checkpoint molecules represent a partial exception: they are *molecular* signals (cheap to produce) that *mask* the dynamical signal (expensive to measure). A cancer cell expressing PD-L1 is not producing high-D dynamics; it is interfering with the immune cell's ability to measure dynamics accurately.

This is why checkpoint molecules are evolutionary innovations: they provide cheap camouflage for an otherwise honest signal. But they are not perfect—checkpoint blockade removes the mask and reveals the underlying signal (Figure 3).

### 4.4 Implications for Immune Evasion

This framework predicts a hierarchy of evasion strategies:

1. **No evasion:** Cancer detected early and cleared. Most initiated tumors.
2. **Cheap molecular camouflage:** Checkpoint expression, antigen loss. Works temporarily; defeated by blockade or antigen-independent surveillance.
3. **Costly dynamical mimicry:** Actually investing in complexity. Works, but cell loses replicative advantage—effectively becoming a cooperator again.
4. **Threshold manipulation:** Inducing local immunosuppression (Tregs, MDSCs) to raise detection threshold. Does not fake the signal; changes the decision rule.

Only strategies 2 and 4 allow a cell to remain a defector while evading detection. Both represent “moving the goalposts” rather than faking the signal. Checkpoint blockade defeats strategy 2; strategies targeting the tumor microenvironment address strategy 4.

### 4.5 The Mutation Burden Paradox

A puzzle in immuno-oncology: tumor mutation burden (TMB) is used as a biomarker for checkpoint blockade response, on the theory that more mutations create more neoantigens

for T cells to recognize. Yet the correlation is weak and inconsistent:

- Some high-TMB tumors are poorly immunogenic and do not respond to checkpoint blockade.
- Some low-TMB tumors are highly immunogenic and respond well.
- Within tumor types, TMB explains only a fraction of response variance.

The dimensional framework resolves this paradox. Immunogenicity is not about mutation count; it is about *attractor structure*. A tumor with many mutations but a coherent replicative program (stable low-D attractor) may be harder to detect than a tumor with few mutations but chaotic, unstable dynamics (fluctuating dimensionality that periodically dips below threshold).

TMB may correlate with response not because mutations create recognizable antigens, but because genomic instability destabilizes the tumor attractor, making it harder to maintain consistent dimensional camouflage. The immune system is not “reading” neoantigens—it is detecting dynamical instability.

This predicts that transcriptional heterogeneity, attractor stability metrics, and single-cell dimensionality measures should outperform TMB as biomarkers for immunotherapy response.

## 4.6 Dynamical Friction and Immune Exhaustion

A high-dimensional immune cell is not isolated. Surveillance requires physical coupling to targets—through receptor engagement, cytoskeletal contacts, and shared biochemical fields. In our framework, these interactions are not neutral: they generate **dimensional friction**.

Consider an immune cell with effective dimensionality  $D_{\text{immune}}$  probing a target of dimensionality  $D_{\text{target}}$ . When the target is healthy tissue,  $D_{\text{target}} \approx D_{\text{immune}}$ : the two systems inhabit similar high-dimensional attractors, and synchronization probes can be performed

with little distortion of the immune cell's own dynamics. When the target is cancerous or virally hijacked,  $D_{\text{target}} \ll D_{\text{immune}}$ : the target occupies a much simpler attractor, with only a few dominant modes. Coupling the two pulls the joint system toward that simpler manifold.

To remain in a high-dimensional regime while coupled to a low-dimensional target, the immune cell must continuously pump energy into the modes that the interaction tends to damp. This generates an effective friction cost:

$$C_{\text{friction}} \propto k (D_{\text{immune}} - D_{\text{target}})^2 \quad (7)$$

where  $k$  is the coupling strength (surveillance intensity). When  $D_{\text{target}} \approx D_{\text{immune}}$  the cost is negligible; when  $D_{\text{target}} \ll D_{\text{immune}}$  the cost is maximal.

Let  $E(t)$  denote the metabolic reserve available for maintaining high-dimensional dynamics, and  $P_{\text{met}}$  the rate at which the organism can resupply this reserve. On the timescale of a sustained interaction:

$$\frac{dE}{dt} = P_{\text{met}} - C_0 - k (D_{\text{immune}} - D_{\text{target}})^2 \quad (8)$$

where  $C_0$  is the baseline cost of being an active immune cell. If resupply is limited ( $P_{\text{met}} \approx C_0$ ), the reserve decays and the **exhaustion time** scales as:

$$t_{\text{exhaust}} \sim \frac{E_0}{k (D_{\text{immune}} - D_{\text{target}})^2} \quad (9)$$

Deeply collapsed, strongly coupled tumors thus exhaust high-dimensional surveillance much more rapidly than healthy tissue does.

Dimensional friction also acts directly on the immune cell's complexity. In the absence of active maintenance, coupling to a low- $D$  target drives  $D_{\text{immune}}$  toward  $D_{\text{target}}$ . A simple

relaxation model captures this competition:

$$\frac{dD_{\text{immune}}}{dt} = -\gamma (D_{\text{immune}} - D_{\text{target}}) + \eta P_{\text{met}} \quad (10)$$

The first term is dimensional friction: in absence of energy input, you relax exponentially toward the target’s dimensionality. The second term is active complexity pumping: metabolism pushes you back up the manifold. In steady state, the **maximum sustainable dimensional gap** is:

$$D_{\text{immune}}^* - D_{\text{target}} = \frac{\eta}{\gamma} P_{\text{met}} \quad (11)$$

This equation is central. With aging and chronic stress, organism-level loss of complexity [11] reduces  $P_{\text{met}}$ , shrinking the dimensional gap the immune system can sustain. High-dimensional surveillance becomes a luxury the organism can no longer afford. The observed phenomenon of **T-cell exhaustion** can thus be interpreted as forced dimensional collapse: immune cells are dragged into low-dimensional attractor basins by repeated coupling to collapsed targets under an insufficient complexity budget.

This is not merely “tiredness”—it is a thermodynamic phase transition. The immune cell literally cannot maintain enough dimensions to distinguish self from non-self. It has been pulled into the tumor’s attractor.

## 5 Inflammation as Controlled Destabilization

### 5.1 The Function of Inflammation

Inflammation is evolutionarily ancient and highly conserved. If it simply caused tissue damage and enabled cancer, it would have been selected against. Its persistence implies function.

We propose that inflammation is **controlled destabilization**: a transient increase in tissue volatility that enables cellular state transitions. When defection is detected, the optimal response is often not direct killing but facilitated transition:

- **Apoptosis:** The defector transitions to death.
- **Clearance:** Phagocytes remove debris.
- **Regeneration:** Stem cells replace lost tissue.
- **Remodeling:** Tissue architecture reorganizes.

All of these require state transitions. Inflammation “shakes” the attractor landscape, lowering barriers between states and enabling transitions that would otherwise be kinetically trapped. This aligns with the “critical transition” framework in aging biology [11], where loss of resilience (shallowing of attractor basins) precedes abrupt system state changes. Inflammation modulates this resilience actively: acute inflammation transiently destabilizes to enable clearance; chronic inflammation permanently erodes the landscape.

In dynamical terms, inflammation increases the effective temperature of the tissue:

$$P(\text{transition}) \propto \exp\left(-\frac{\Delta E}{k_B T_{\text{eff}}}\right) \quad (12)$$

where  $\Delta E$  is the barrier height between attractors and  $T_{\text{eff}}$  is an effective temperature set by inflammatory signals. Inflammation raises  $T_{\text{eff}}$ , increasing transition probability.

## 5.2 Acute vs. Chronic Inflammation

Acute inflammation is adaptive: destabilize, clear the problem, restabilize. The tissue returns to a high-dimensional attractor with defectors removed.

Chronic inflammation is maladaptive: persistent destabilization without resolution. The tissue remains in a high- $T_{\text{eff}}$  state, where:

1. Attractor basins are shallow (cells easily perturbed).
2. State transitions are frequent (including transitions to defection).
3. Immune thresholds are raised (tolerance under persistent alarm).

4. Defectors accumulate (production exceeds clearance).

This is the mechanism linking chronic inflammation to cancer: not “inflammation causes mutations” but “inflammation destabilizes the attractor landscape, enabling escape to low-D states while simultaneously raising detection thresholds.”

## 6 The HPA Axis as Threshold Modulator

### 6.1 Cortisol and Immune Tolerance

The hypothalamic-pituitary-adrenal (HPA) axis regulates the stress response, with cortisol as its primary effector. Cortisol is immunosuppressive: it reduces inflammatory signaling, suppresses lymphocyte proliferation, and raises the threshold for immune activation.

In our framework, cortisol raises the dimensional detection threshold: under stress, tolerate more deviation from high-D tissue norms. This makes evolutionary sense. During acute stress (predator attack, injury, starvation), the organism cannot afford to divert resources to immune surveillance. Defectors that would normally be eliminated are temporarily tolerated.

The problem is chronic stress. Persistently elevated cortisol produces persistently raised thresholds. Meanwhile, chronic stress often co-occurs with other destabilizing factors (poor nutrition, sleep disruption, social isolation) that shallow the attractor landscape. The combination is toxic: more cells falling into low-D states, fewer being caught.

### 6.2 Social Buffering and the Cooperative Niche

Social support buffers HPA-axis reactivity [3]. Individuals embedded in cooperative social networks show reduced cortisol responses to stressors. This has downstream effects on immune function: lower chronic cortisol means lower detection thresholds means better surveillance.

This connects to Sierra et al.’s [4] finding that cooperative species have lower cancer rates. Cooperation operates at multiple scales:

1. **Cellular**: Immune enforcement against defecting cells.
2. **Organismal**: Developmental coherence that stabilizes tissue attractors.
3. **Social**: Group buffering that reduces HPA activation.

The same dynamical structure—attractor stabilization via cooperative interaction—appears at each scale. This is “fractal cooperation”: the mechanism is scale-invariant.

### 6.3 The Entropic Cascade: When the Sensor Fails

Crucially, the immune system is itself a physiological system subject to the “loss of structural and functional complexity” described by Cohen et al. [11]. As organismal metabolism declines with age, the high energetic cost of maintaining high-dimensional dynamics becomes unsustainable. This creates thermodynamic pressure to shed complexity.

This leads to a double failure mode:

1. **Target generation**: Tissue cells fall into low-dimensional attractors (cancer/senescence) to save energy.
2. **Sensor blindness**: Immune cells simultaneously lose the dynamical bandwidth required to detect these states.

The result is an **entropic cascade**: loss of complexity in one module (e.g., mitochondrial function) propagates through the network, forcing simplification of the surveillance system, which in turn permits survival of simpler, defecting cells. The organism cannot afford both complex tissue dynamics *and* complex surveillance dynamics; something must give.

This explains the “blurry lens” problem of aging immunity. A low- $D_{\text{eff}}$  sensor cannot reliably distinguish a high- $D_{\text{eff}}$  healthy cell from a low- $D_{\text{eff}}$  cancer cell—the measuring instrument has lost resolution. Immunosenescence is not merely quantitative decline (fewer T cells, slower proliferation); it is qualitative degradation of the measurement apparatus itself.

The thermodynamic trap is clear: maintaining complexity requires energy, but aging reduces energy availability. The organism faces a choice between complex tissue function and complex surveillance. Neither can be sustained indefinitely. Cancer incidence rises with age not only because mutations accumulate, but because the surveillance system can no longer afford to maintain the dimensional bandwidth necessary for detection.

## 6.4 Empirical Evidence: Cancer Accelerates Biological Aging

If the entropic cascade is real, cancer should accelerate biological aging—not just because of treatment toxicity, but because the tumor acts as a low-dimensional sink draining complexity from the organism. This prediction is strongly supported by clinical evidence.

Cancer survivors show dramatic acceleration of biological aging across multiple metrics [12, 13]:

- **Epigenetic clocks:** Survivors are 5–16 years biologically older than age-matched controls; the pace of aging is approximately 19% faster in survivors versus comparators.
- **Cellular senescence:** Frail cancer survivors show p16<sup>INK4a</sup> expression levels representing a 25-year age acceleration compared to robust survivors.
- **Telomere attrition:** Chemotherapy and radiation accelerate telomere shortening beyond normal aging trajectories.
- **Chronic inflammation:** IL-6 and other inflammatory markers remain chronically elevated in survivors, consistent with persistent attractor destabilization.

- **Frailty:** Cancer survivors exhibit frailty phenotypes typically seen in individuals decades older.

Critically, this acceleration occurs even in young adult survivors of childhood cancers—ruling out explanations based solely on pre-existing age-related decline. The entropic cascade model predicts this: cancer is not merely a disease that kills; it is a *dimensional drain* that ages the entire organism by forcing sustained high-cost surveillance against a low-complexity adversary. The immune system hemorrhages complexity fighting the tumor; the tumor wins even when it loses, by accelerating organismal aging.

This reframes cancer survivorship. “Curing” the cancer does not restore the lost complexity—the entropic debt has been paid. Interventions targeting biological age may be as important as tumor-directed therapy for long-term survivor outcomes.

## 6.5 Complexity Restoration: The Mechanistic Basis of Lifestyle Medicine

If cancer drains complexity, recovery requires *complexity restoration*. The standard “lifestyle medicine” recommendations—exercise, diet, social support, stress reduction—are not vague wellness advice. Each directly counteracts the entropic cascade:

- **Exercise:** Stimulates mitochondrial biogenesis, increasing ATP production capacity.

More energy means the organism can afford to maintain high- $D_{\text{eff}}$  dynamics. Meta-analyses show exercise reduces pro-inflammatory markers (standardized mean difference  $-0.2$ ), increases NK cell and cytotoxic T cell counts, and inhibits immunosenescence [14]. Exercise literally reverses the dimensional collapse.

But exercise does something deeper than provide energy: it *synchronizes* the organism. During physical activity, the cardiovascular, respiratory, muscular, and metabolic systems enter a coordinated state—a coherent, goal-directed reduction in dimensionality. This is not pathological collapse; it is purposeful alignment. And this alignment

may enhance detection: when the entire tissue environment is synchronized toward a common attractor (movement, exertion), a cancer cell running its own autonomous program becomes *more visible*, not less. It is the one cell that cannot join the march. Exercise may function as a “contrast agent” for immune surveillance, reducing background noise and highlighting defectors through their failure to synchronize with the organismal program.

- **Social support:** Buffers HPA-axis reactivity, reducing chronic cortisol. Lower cortisol means lower detection thresholds and deeper attractor basins. Cancer patients with high-quality social support show higher NK cell activity and lower cortisol; social network size reduces mortality risk by 20% [15, 16].
- **Stress management:** Modulates the neuroendocrine-immune axis. Mindfulness, CBT, and relaxation techniques alter immune parameters in ways that may influence disease progression. Flatter diurnal cortisol slopes predict shorter survival—stress management restores normal cortisol dynamics.
- **Purpose and meaning:** This is the deepest intervention. Exercise and social support don’t just provide *energy* for complexity—they provide *structure*. Goals, relationships, and meaningful activity create the attractor landscape into which complexity is organized. Without purpose, high dimensionality is mere noise (pathological, like seizures). With purpose, high dimensionality becomes coherent function.

The distinction matters. A cell can have high-dimensional dynamics that are *uncoordinated*—fluctuating wildly without serving any function. What healthy tissue exhibits is high-dimensional dynamics that are *embedded in functional attractors*: complex, but purposefully complex. The organism needs both: sufficient metabolic capacity to maintain complexity, and sufficient structure (goals, relationships, meaning) to organize that complexity into function.

This explains why “having something to live for” predicts cancer survival beyond what medical variables explain. Purpose is not mysticism—it is attractor architecture. A person with goals and relationships has deeper, more stable attractor basins. Their complexity has somewhere to go. A person without purpose has shallower basins; their dynamics are more easily destabilized; they are more vulnerable to the entropic cascade.

Cancer recovery, on this view, is not merely about eliminating tumor cells. It is about rebuilding the dimensional capacity and attractor structure that the cancer eroded. This is a long-term project requiring sustained investment in the systems—metabolic, social, psychological—that generate and organize complexity. The tumor may be gone, but the landscape must be restored.

## 7 Price Equation Formalization

### 7.1 Within-Organism Selection

Let  $z_i$  be the replicative rate of cell  $i$  in an organism, and  $w_i$  be the cell’s fitness (survival probability  $\times$  replication rate). In the absence of immune enforcement:

$$\Delta \bar{z} = \text{Cov}(w_i, z_i) > 0 \quad (13)$$

Cells with higher replicative rates have higher fitness; the population evolves toward increased replication (cancer).

Immune enforcement adds a negative term: high- $z$  cells are preferentially eliminated. If detection is based on dimensionality, and high- $z$  cells have low  $D_{\text{eff}}$ :

$$w_i = w_0 \cdot r(z_i) \cdot s(D_{\text{eff}}(z_i), \theta) \quad (14)$$

where  $r(z)$  is the replicative advantage of rate  $z$  and  $s(D_{\text{eff}}, \theta)$  is the survival probability given dimensionality  $D_{\text{eff}}$  and detection threshold  $\theta$ . If  $s$  decreases steeply as  $D_{\text{eff}}$  falls below

threshold:

$$\Delta \bar{z} = \text{Cov}(w_0 \cdot r \cdot s, z) \approx 0 \quad (15)$$

when enforcement is effective. The replicative advantage is balanced by immune elimination.

## 7.2 Threshold Modulation

The threshold  $\theta$  is modulated by organismal state:

$$\theta = \theta_0 + \alpha \cdot C \quad (16)$$

where  $C$  is cortisol (or a proxy for HPA activation) and  $\alpha > 0$ . Under stress,  $\theta$  rises, and the survival function  $s(D_{\text{eff}}, \theta)$  becomes more permissive. Low-D cells that would normally be eliminated now survive.

If stress is chronic, the population evolves: cells with lower  $D_{\text{eff}}$  accumulate because  $s$  remains high. This is the Price-equation formalization of “chronic stress enables cancer.”

# 8 Evolutionary Elaboration of Immunity

## 8.1 From Simple Surveillance to Adaptive Immunity

Early multicellular organisms likely had simple dimensional surveillance: phagocytes that engulfed cells falling below a complexity threshold. This is essentially what macrophages still do.

The evolution of adaptive immunity (B cells, T cells, MHC, antibodies) elaborates this basic mechanism:

1. **TCR/BCR diversity:** More sophisticated synchronization probes. Instead of a single

coupling measurement, the adaptive immune system generates diverse probes that sample the target cell's response space.

2. **MHC presentation:** A window into the target cell's internal state. Peptide-MHC complexes report what proteins the cell is making, providing additional dimensionality information beyond surface coupling.
3. **Clonal selection:** Probes that detect low-D signatures are amplified; probes that respond to high-D (self) signatures are eliminated (negative selection) or suppressed (tolerance).
4. **Memory:** Successful probe configurations are stored, enabling rapid response to recurring threats.

The underlying principle remains dimensional surveillance, but the implementation becomes more sophisticated.

## 8.2 The Autoimmunity-Cancer Tradeoff

If detection thresholds are set too low (aggressive surveillance), the system attacks normal cells that transiently dip in dimensionality. This is autoimmunity.

If thresholds are set too high (permissive surveillance), the system tolerates low-D cells that should be eliminated. This is cancer susceptibility.

Evolution has tuned the threshold to balance these costs. But the optimal threshold depends on environment:

- High pathogen load: Lower threshold beneficial (catch infections early, tolerate autoimmune cost).
- Low pathogen load: Higher threshold beneficial (avoid autoimmunity, tolerate cancer cost).

Modern humans live in low-pathogen environments but carry immune systems tuned for high-pathogen ancestral conditions. This may explain rising autoimmunity rates.

### 8.3 Autoimmunity as Sensor Collapse: The Blunt Instrument Problem

The threshold model captures part of autoimmunity, but a deeper mechanism emerges from the dynamical friction analysis: **autoimmunity can result from dimensional collapse in the sensor itself.**

A high-dimensional immune cell possesses sufficient degrees of freedom to distinguish “Healthy Self” (high-D, stable) from “Stressed Self” (high-D but shifted in state space) from “Tumor” (low-D, collapsed). It has high *discriminatory resolution*. However, if the immune cell itself suffers dimensional collapse—due to dynamical friction from chronic antigen exposure, persistent inflammation, or organismal aging—it loses this resolution.

Formally, discrimination requires the sensor’s dimensionality to exceed the manifold dimension separating target states. When  $D_{\text{sensor}} \ll D_{\text{required}}$ , distinct target states *alias* onto the same low-dimensional projection. The sensor cannot distinguish them.

This forces the collapsed sensor into binary defaults:

- **Indiscriminate Attack (Autoimmunity):** The collapsed sensor defaults to a rigid inflammatory program, firing on normal tissues because it lacks the dimensional bandwidth to process inhibitory context signals.
- **Indiscriminate Tolerance (Exhaustion):** The sensor defaults to quiescence, ignoring actual threats because discrimination is metabolically impossible.

This explains the paradox of *inflamm-aging*: why do elderly individuals exhibit weak immunity against novel pathogens yet strong autoimmune inflammation? Because their immune systems have collapsed into low-dimensional attractors that fire indiscriminately. The aging immune system is simultaneously ineffective (low sensitivity to genuine threats)

and non-specific (high false-positive rate against self) because it has lost the complexity required for precision.

Autoimmunity, on this view, is not merely a software error (wrong target recognized) but a **hardware failure** (broken measurement apparatus). The “lens” has degraded; the image is blurred.

## 8.4 Systemic Hyper-Coupling: The Autism-Autoimmunity Connection

The logic of dimensional collapse extends beyond immunity. Consider autism, characterized by excessive neural connectivity and phase-locking. Paradoxically, hyper-connectivity *reduces* neural complexity: when coupling strength  $K$  exceeds a critical threshold  $K_c$ , the system undergoes a phase transition to global synchronization. Effective dimensionality collapses toward unity as the entire network beats in unison.

This suggests a unified etiology for the observed correlation between autism and autoimmunity: a systemic bias toward **hyper-coupling**. An organism with a genetic or developmental set-point for high coupling strength will exhibit dimensional collapse in every complex adaptive system:

- **Neural Hyper-sync (Autism):** Excessive phase-locking reduces cognitive flexibility.

The brain cannot maintain the “chimera states” (partial synchronization) required for complex computation. It gets stuck in deep, repetitive attractors (stereotypy) and is overwhelmed by sensory input because it cannot dampen the synchronization cascade.

- **Immune Hyper-sync (Autoimmunity):** Excessive receptor sensitivity or signaling gain causes the immune sensor to collapse into “Alarm” states too easily. The system phase-locks to *any* input, losing the resolution to distinguish self from non-self.

Both are manifestations of a system that has lost the ability to maintain the *critical* regime between order and chaos, collapsing instead into excessive order (low dimensionality).

Autism and autoimmunity are not separate diseases but the **same dynamical parameter error** ( $K > K_c$ ) expressing itself in two different substrates.

This has therapeutic implications: interventions that reduce coupling gain or restore criticality might benefit both conditions. It also reinforces the central thesis: **health is high dimensionality maintained at criticality; disease is dimensional collapse toward excessive order or chaos.**

## 9 Discussion

### 9.1 Predictions

The dimensional surveillance hypothesis makes several testable predictions:

1. **Dynamical complexity correlates with immune evasion.** Tumors that evade immunity should show higher apparent dimensionality (either genuine or camouflaged) than tumors that are cleared.
2. **Checkpoint blockade reveals dimensionality.** Pre- and post-treatment comparisons should show that responding tumors had low-D dynamics masked by checkpoint signaling, while non-responders had genuinely high-D dynamics (or other evasion mechanisms).
3. **Chronic stress impairs dimensional detection.** Immunological assays under chronic cortisol exposure should show reduced discrimination between high-D and low-D targets.
4. **Social buffering restores detection.** Individuals with strong social support should show better immune discrimination, measurable via functional assays.
5. **Exercise enhances surveillance.** Consistent with existing literature: exercise reduces cortisol, stabilizes attractors, and should improve dimensional detection.

## 9.2 Relation to Existing Frameworks

Our proposal complements rather than replaces existing immunology:

- **Antigen recognition** remains important, especially for infection. But dimensional surveillance provides the deeper substrate.
- **Danger signals** (DAMPs) are downstream of dimensional collapse: damaged cells release DAMPs because their dynamics have destabilized.
- **Immunoediting** (elimination, equilibrium, escape) maps onto our framework: elimination = low-D cells cleared; equilibrium = marginal cells at threshold; escape = cells that have acquired high-D camouflage.

## 9.3 The Immune Synapse as a Dimensional Bottleneck

Biological systems must constantly bridge two regimes: the continuous, high-dimensional regime of physical dynamics (protein vibrations, ion fluxes, metabolic oscillations) and the discrete, low-dimensional regime of biological control (mitosis, apoptosis, differentiation).

The immune synapse represents the ultimate dimensional bottleneck. The input is the “analog” dynamical coupling between immune cell and target—a high-dimensional, continuous-time interaction involving thousands of molecular degrees of freedom. The output, however, must be digital: the cytotoxic granule release mechanism is binary. It fires or it does not.

In this view, the immune system is not recognizing pre-existing labels; it is **generating codes**. “Self” and “Non-self” are not intrinsic properties of cells; they are the compressed symbolic outputs generated when high-dimensional dynamics are forced through the kill/no-kill bottleneck. The immune synapse functions as an Identification Friend or Foe (IFF) transponder system, but one where the “squawk code” is derived in real-time from the dynamical stability of the target.

This explains why the system is fragile to noise (autoimmunity) and spoofing (cancer). The codeforming process is lossy. When a high-dimensional state (healthy tissue) is compressed into a 1-bit code (Tolerate), massive amounts of information are discarded. If dimensional friction prevents the sensor from fully exploring the target’s state space, the compression fails, and the wrong code is generated. T-cell exhaustion is effectively a failure of this coding machinery—the bandwidth required to generate the correct IFF symbol exceeds the metabolic capacity of the bottleneck.

This framing resolves the famous “combinatorial explosion” paradox: how can  $\sim 10^8$  receptors cover  $\sim 10^{15}$  possible antigens? The answer is that receptors are not 1-to-1 keys for locks; they are **basis vectors** that project the high-dimensional target space onto a low-dimensional manifold of coupling strengths. Antibodies bind to generic dynamical classes (molecular motifs, vibrational modes), not specific molecular identities. Cross-reactivity is not a bug but the essential feature: a single receptor covers vast swathes of “low-D defector” space because low-dimensionality itself is the conserved signal.

## 9.4 Comparative Dynamics: The Regeneration-Cancer Tradeoff

Our framework resolves the long-standing paradox of regeneration versus cancer: why do highly regenerative organisms (axolotls, planaria) show extreme cancer resistance, while mammals exhibit limited regeneration but high cancer susceptibility?

**The Mammalian Strategy (High-D / Deep Attractors):** Mammals enforce cooperation by locking cells into deep, high-dimensional attractor basins. This creates “high friction” against state changes.

- **Benefit:** Extreme resistance to casual defection. Small perturbations cannot push a cell out of its deep well.
- **Cost:** Loss of regenerative capacity. The same walls that prevent cancer also prevent dedifferentiation for healing. When cancer does occur (jailbreak), it implies the cell has

overcome a massive barrier, resulting in aggressive, hard-to-control phenotypes with high intratumor heterogeneity.

**The Axolotl Strategy (Low-D / Shallow Attractors):** Regenerative species maintain shallow, plastic attractor basins.

- **Benefit:** High regenerative capacity. Cells can traverse between “differentiated” and “stem-like” states with minimal energy cost.
- **Cost:** High theoretical instability.
- **The Solution:** Instead of *preventing* movement (walls), these systems rely on *active guidance*. As described by Levin [17], these organisms maintain robust bioelectric networks that continuously enforce the “target morphology.” They are masters of dimensional shepherding: they do not fear cancer because their physiology is built to manage dynamic state transitions, actively guiding wayward cells back to the tissue attractor rather than simply killing them.

Human cancer vulnerability is thus the “price of permanence”—the cost of relying on rigid structural constraints rather than the active bioelectric management seen in regenerative species. Scarring is a high-D, panic-mode response to injury: lock everything down to prevent dimensional collapse.

This suggests a dimensional phase diagram across species: regeneration capacity, cancer incidence, and effective dimensionality co-vary. Higher-D organisms have less regeneration and fewer “simple” cancers, but when cancer escapes, it is high-D cancer—heterogeneous, plastic, and evasive. Lower-D organisms with active guidance suppress even simple cancers and can re-run developmental programs to clear aberrant cells.

## 9.5 Implications for Therapy

If immunity is dimensional surveillance, therapeutic strategies should aim to:

1. **Reveal true dimensionality:** Checkpoint blockade does this by removing camouflage.
2. **Force dimensional collapse:** Therapies that push tumor cells into obviously low-D states (differentiation therapy, metabolic disruption) should enhance immune recognition.
3. **Stabilize tissue attractors:** Systemic interventions that deepen attractor basins (exercise, stress reduction, anti-inflammatory lifestyle) should reduce the supply of low-D cells.
4. **Restore detection thresholds:** Interventions that normalize HPA function should improve surveillance.

This provides a unified framework for understanding why diverse interventions (checkpoint blockade, differentiation therapy, exercise, stress reduction) all affect cancer outcomes.

## 9.6 Why Immunotherapy Is Reserved for Advanced Disease

A striking clinical pattern deserves attention: immunotherapy (checkpoint blockade, CAR-T) is typically reserved for patients with advanced, metastatic disease—the sickest patients. From a dimensional surveillance perspective, this may not be coincidental.

Advanced cancer patients typically exhibit:

- High tumor burden → systemic inflammation
- Chronic stress → elevated cortisol
- Cachexia → metabolic dysregulation
- Treatment history → accumulated tissue damage

All of these factors produce **shallow attractor basins** throughout the organism. In our framework, this has competing implications:

1. **Easier state transitions:** Shallow basins mean cells can be pushed into new states more easily—including death. This may enhance immunotherapy efficacy.
2. **More side effects:** Shallow basins also mean normal cells are more likely to transiently dip below detection thresholds, triggering autoimmunity. Checkpoint blockade in this context is particularly prone to immune-related adverse events.
3. **Less durable responses:** Unstable attractors don't stay put. Even if immunotherapy clears a tumor, the destabilized landscape may permit rapid recurrence or new malignancies.

This suggests that immunotherapy might work differently—potentially better—if applied earlier, when attractor landscapes are more stable. However, clinical practice reserves it for late-stage disease due to toxicity concerns and cost. The dimensional framework suggests this may be suboptimal.

## 9.7 Limitations: Measuring Effective Dimensionality

A fundamental limitation of this framework is that effective dimensionality ( $D_{\text{eff}}$ ) is a theoretical construct that cannot be directly measured *in vivo*. In our simulations, we compute  $D_{\text{eff}}$  via principal component analysis of oscillator trajectories. In living cells, we cannot observe the full state space.

What we *can* measure are proxies:

- **Gene expression dimensionality:** Single-cell RNA sequencing reveals transcriptional state space. Cancer cells often show reduced transcriptional diversity.
- **Metabolic complexity:** Metabolomic profiling can capture metabolic network activity patterns.
- **Calcium dynamics:** Imaging of intracellular calcium reveals oscillatory complexity that may correlate with  $D_{\text{eff}}$ .

- **Electrophysiology:** For excitable cells, action potential patterns provide a window into dynamical complexity.

However, none of these is  $D_{\text{eff}}$  itself. The theory is testable in principle—high-D cells should evade detection less than low-D cells, controlling for other factors—but direct measurement of the proposed signal remains a challenge.

We do not claim to know the order of magnitude of  $D_{\text{eff}}$  for real cells. In simulation, we use  $D_{\text{eff}} \sim 10^1$ ; in reality, cellular state spaces may have  $D_{\text{eff}} \sim 10^2$  or higher. The key claim is not about absolute values but about the *difference* between cooperating and defecting cells: defectors should be measurably lower in whatever complexity metric is used.

## 10 Conclusion

The immune system evolved to solve a cooperation problem: detecting and eliminating cells that defect from the multicellular program. The signature of defection is dynamical collapse—reduced effective dimensionality reflecting escape from complex tissue attractors into simple replicative or dysfunctional states.

Immune receptors function as synchronization probes that measure target cell dimensionality. Inflammation is controlled destabilization that facilitates state transitions. The HPA axis modulates detection thresholds based on organismal state.

This framework unifies cancer immunology, autoimmunity, and chronic inflammation under a single principle: immunity is cooperation enforcement via dimensional surveillance. The Price equation provides the formal structure: within-organism selection favors defection; immune enforcement suppresses it; threshold modulation trades off surveillance costs against defection costs.

Cancer, autoimmunity, and chronic inflammatory disease are not separate problems—they are different failure modes of a single system. Understanding immunity as dimensional surveillance may enable more principled therapeutic strategies.

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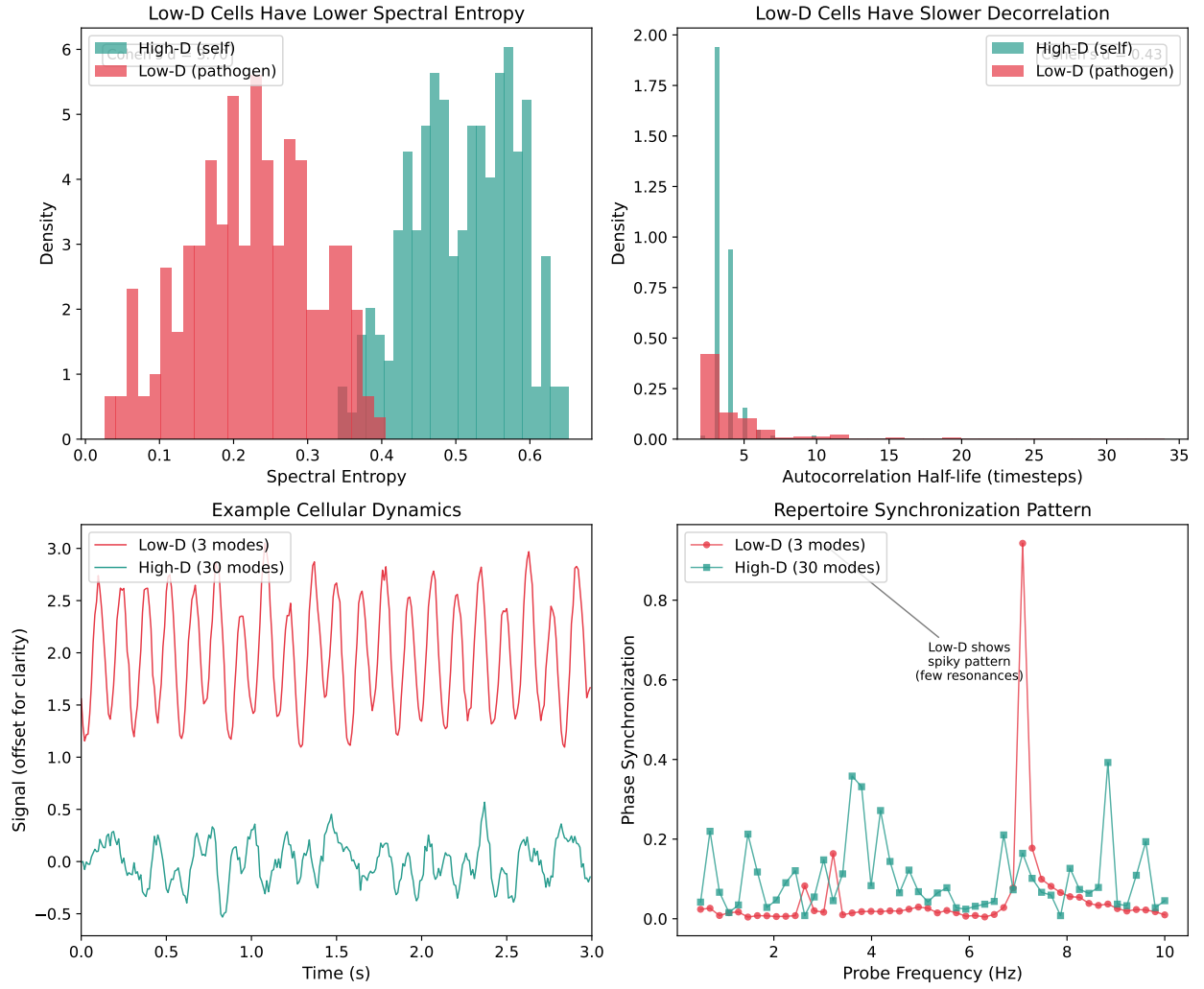


Figure 1: Temporal dynamics discriminate dimensionality. (A) Spectral entropy distributions show near-complete separation between low-D (pathogen-like) and high-D (self-like) dynamics. (B) Autocorrelation decay is faster for high-D signals. (C) Example time series: low-D (3 modes) shows coherent oscillation; high-D (30 modes) shows complex dynamics. (D) Synchronization pattern across probe frequencies: low-D shows sharp resonance peaks at its few active frequencies; high-D shows uniform low-level response.

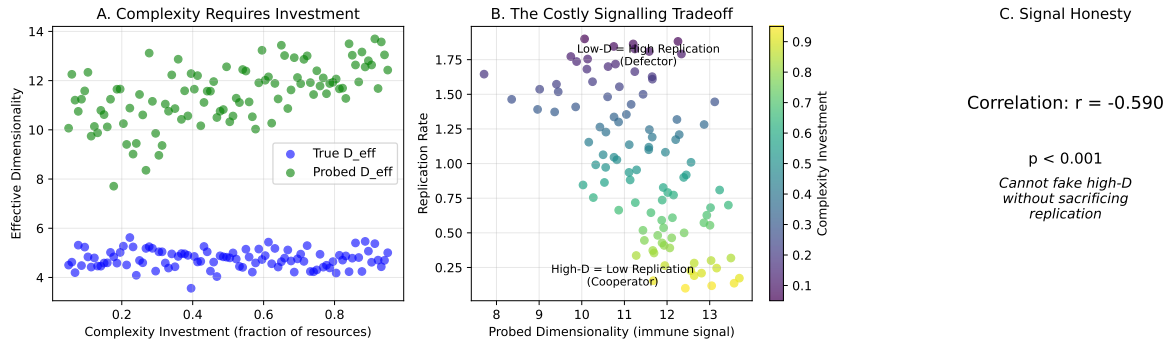


Figure 2: The costly signalling tradeoff. (A) Effective dimensionality requires metabolic investment. (B) High replication rate correlates with low probed dimensionality ( $r = -0.59$ ). Defectors (low-D, high replication) cluster in the upper left; cooperators (high-D, low replication) in the lower right. (C) This negative correlation ensures dimensional complexity is an honest signal.

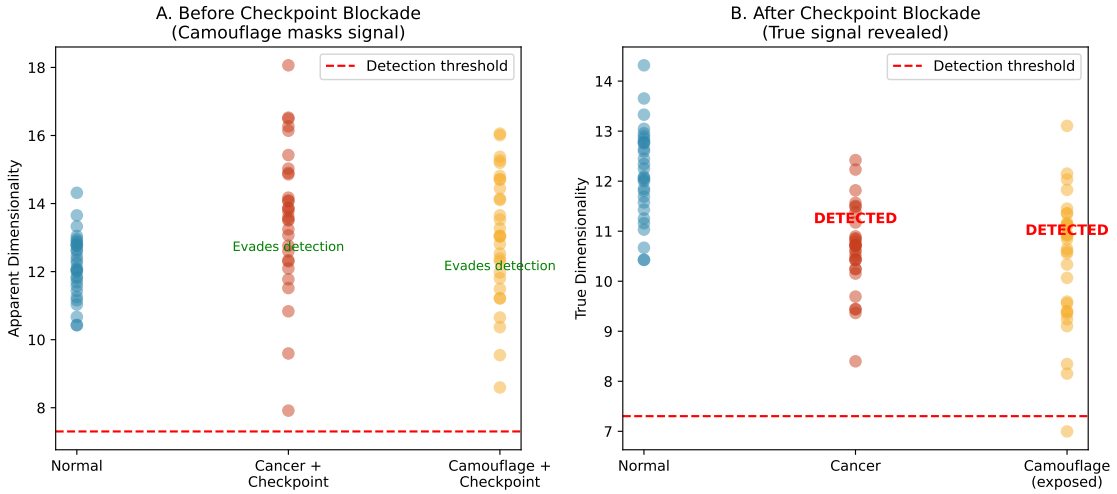


Figure 3: **Checkpoint Blockade Reveals True Dimensionality.** (A) With checkpoint protection, cancer cells appear to have higher dimensionality (camouflage works). (B) After checkpoint blockade, the true low-D signal is exposed and cells are detected.

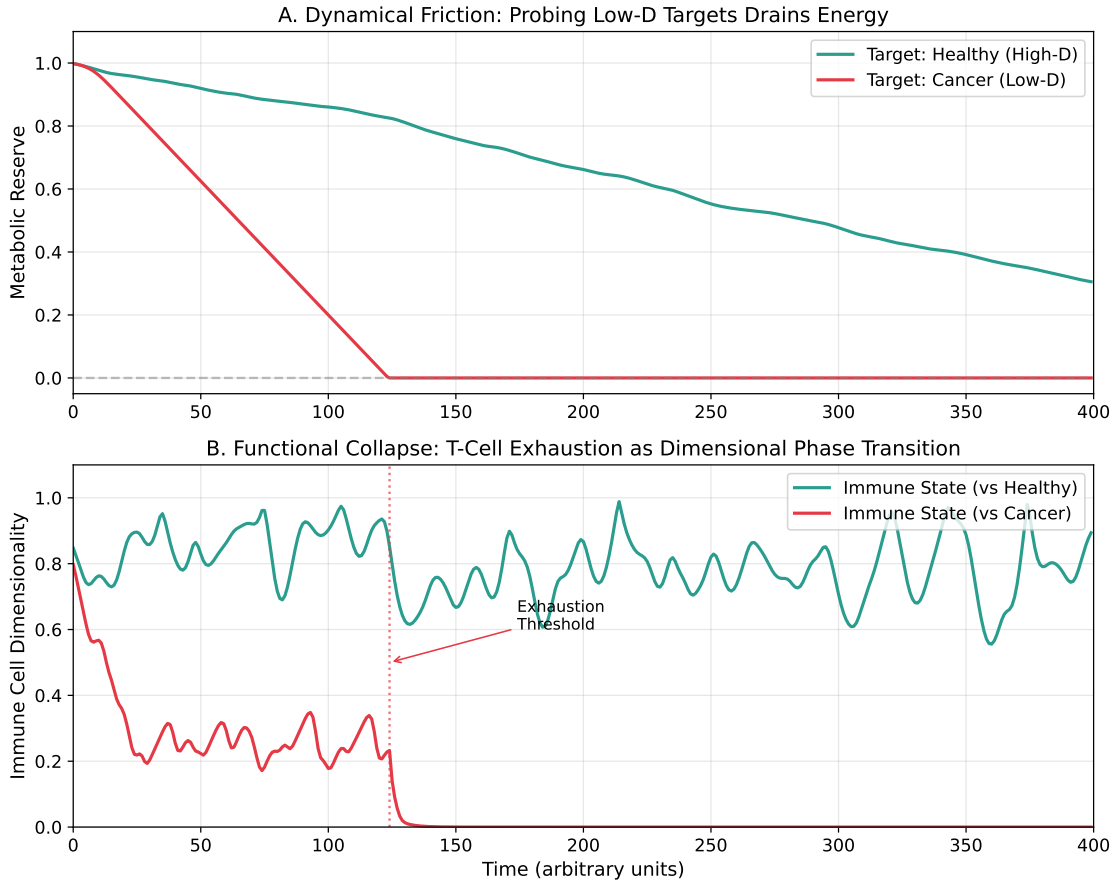


Figure 4: **Dynamical Friction and Exhaustion.** (A) Probing a low-dimensional target (red) drains metabolic reserve rapidly compared to a high-dimensional target (teal) due to the cost of resisting synchronization. (B) When reserve is depleted, the immune cell undergoes a phase transition, collapsing into a low-dimensional exhausted state. Simulation uses coupled Kuramoto oscillators where high-D corresponds to desynchronized dynamics and low-D corresponds to synchronized dynamics. See Equation 9 for the scaling relationship.