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**Adaptive Regenerative Error Due to Loss of Cellular Reference Pattern: A Hypothesis of Dominant Substitution in Chronic Inflammatory Microenvironments**

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# Adaptive Regenerative Error Due to Loss of Cellular Reference Pattern: A Hypothesis of Dominant Substitution in Chronic Inflammatory Microenvironments

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## Abstract

Aberrant cellular adaptation is a hallmark of various chronic diseases, including endometriosis, metaplasia, and fibrotic conditions. This paper proposes a novel hypothesis: that such pathological transformations result from a progressive loss of the original cellular reference pattern under sustained inflammatory and dysregulated conditions. Termed the Dominant Substitution Hypothesis, this model suggests that chronic microenvironmental disruption alters regenerative cues, gradually replacing healthy cell phenotypes with adaptive, yet functionally impaired, variants. Once a critical threshold is reached, the adaptive phenotype becomes dominant, perpetuating dysfunction and inhibiting restoration. The hypothesis integrates evidence from tissue plasticity, extracellular matrix disorganization, epigenetic modulation, microbiota-driven signaling, and immune-hormonal imbalance. Implications for diagnosis, prevention, and regenerative therapy are discussed, with a focus on early intervention to preserve cellular identity and interrupt the degenerative cycle.

## Keywords

Regenerative Error, Cellular Pattern Loss, Epigenetic Drift, Phenotypic Substitution, Chronic Inflammation, Tissue Remodeling, Microbiota Signaling, Cellular Plasticity, Endometriosis, Repair Failure Mechanism

## Introduction

Chronic inflammatory conditions often lead to progressive structural and functional changes in tissues, frequently culminating in irreversible degeneration. Despite advancements in understanding cellular plasticity, epigenetics, and immunometabolic

interactions, a unified explanation for the transition from regeneration to dysfunction remains elusive. Current models treat pathological transformations as either genetically driven or immune-mediated, frequently overlooking the role of long-term environmental disruption on cellular identity. Accordingly, this manuscript is presented as a hypothesis-driven conceptual framework rather than an experimental or data-reporting study.

This paper introduces a hypothesis-driven conceptual framework, termed the Dominant Substitution Hypothesis, which proposes that under conditions of persistent tissue stress and unresolved inflammation, the cellular microenvironment becomes progressively dysregulated. This dysregulation disrupts the cues necessary for faithful cellular regeneration, causing new cells to adopt adaptive phenotypes that are not functionally equivalent to the originals. Initially, these adaptations serve as compensatory mechanisms, but over time, their accumulation surpasses a critical threshold. Once the proportion of adaptive cells exceeds that of original phenotypes, the altered state becomes self-perpetuating, locking the tissue into a dysfunctional loop.

This hypothesis builds upon established evidence from regenerative biology, matrix remodeling, epigenetic drift, and gut microbiota signaling. By integrating these domains, it aims to offer a comprehensive model for early identification of degenerative risk and open avenues for interventions focused on preserving or restoring the original regenerative pattern of cellular identity. The purpose of this hypothesis is to provide a unifying explanatory model and to stimulate future experimental and clinical investigation.

## Theoretical Background

To establish the conceptual foundation for the Dominant Substitution Hypothesis, this section reviews the major biological mechanisms that support the plausibility of progressive loss of cellular reference in chronic degenerative conditions.

### Cellular Plasticity and Phenotypic Drift

Cellular plasticity refers to the ability of cells to modify their phenotype in response to environmental stimuli. While essential for development and tissue repair, plasticity can also contribute to pathological transformation under chronic stress. Studies in metaplasia (Slack 2007), epithelial-to-mesenchymal transition (EMT) (Zeisberg and Neilson 2019), and adaptive immune responses demonstrate that cells may adopt noncanonical states when exposed to sustained microenvironmental disruption.

### Extracellular Matrix Disruption and Loss of Structural Cues

The extracellular matrix (ECM) provides not only mechanical support but also biochemical guidance for cell behavior. Chronic inflammation alters ECM composition and stiffness (Wynn and Ramalingam 2012), leading to loss of spatial orientation (Zahir and Weaver 2004), aberrant integrin signaling, and impaired regeneration. Without an

intact ECM scaffold, regenerating cells receive distorted positional information, increasing the risk of phenotypic deviation.

## **Epigenetic Modulation Under Chronic Inflammation**

Epigenetic changes, such as DNA methylation, histone modification, and chromatin remodeling, are responsive to environmental cues and influence gene expression without altering DNA sequence. Persistent inflammatory signals—especially cytokines and oxidative stress—drive epigenetic drift (Feil and Fraga 2012), which may lock cells into maladaptive phenotypes over time (Coussens and Werb 2012).

## **Microbiota-Driven Systemic Signaling**

The intestinal microbiota exerts systemic effects on host tissues (Belkaid and Hand 2014) through metabolites, immune modulation, and hormonal cross-talk. Dysbiosis disrupts these signals, contributing to chronic low-grade inflammation and altering the metabolic and immunological context in which cells regenerate. This adds another layer of environmental complexity that may influence the fidelity of cellular renewal.

## **Threshold Dynamics and Phenotypic Dominance**

Cell populations are shaped by feedback loops and proportional dynamics. When a subset of cells acquires a stable but maladaptive phenotype and their proportion exceeds a critical threshold, they begin to dictate the microenvironmental signals that guide further differentiation. This self-reinforcing shift results in the replacement of original phenotypes by dysfunctional variants, closing the regenerative window.

## **Hypothesis Formulation**

The Dominant Substitution Hypothesis proposes a progressive model of regenerative failure in chronically inflamed tissues. This model rests on the principle that cell regeneration is not only genetically programmed, but also context-dependent, requiring coherent spatial, biochemical, and mechanical cues from the surrounding microenvironment. When these cues become dysregulated over time, the resulting regenerative process is impaired, and new cells increasingly diverge from the original phenotype.

## **Sequential Phases of Adaptive Regeneration**

The hypothesis outlines three core phases:

1. **Initial Dysregulation:** The tissue experiences prolonged inflammation or biochemical imbalance, subtly disrupting cellular guidance signals.

2. Adaptive Regeneration: Regenerating cells begin to adopt partially functional but altered phenotypes, aimed at surviving in a damaged environment.
3. Threshold Substitution: Once the proportion of adaptive cells surpasses a critical threshold, these variants shape the local microenvironment, influencing new cells to conform to the altered state. The original phenotype becomes progressively inaccessible.

## Self-Reinforcing Degeneration

This transition results in a self-perpetuating cycle in which dysfunctional cells dominate, further distorting regenerative signals and stabilizing the maladaptive tissue structure. Rather than representing a random mutation or external aggression, this degenerative process is conceptualized as a systemic loss of reference, where the tissue gradually 'forgets' its original regenerative blueprint.

## Graphical Representation

This process can be modeled as a dynamic threshold function, in which the probability of phenotypic drift increases as the ratio of adapted-to-original cells rises. A tipping point is reached when the majority of environmental signals reflect the adapted phenotype, triggering a shift from reversible adaptation to irreversible substitution. This marks the end of effective tissue recovery and the establishment of chronic pathology.

## Theoretical Case Studies

To illustrate the applicability of the Dominant Substitution Hypothesis across diverse pathological contexts, this section examines several conditions in which maladaptive cellular transformation is a defining feature. These examples are used to demonstrate how a progressive shift in regenerative guidance can underpin chronic dysfunction.

### Endometriosis

Endometriosis involves the presence of endometrial-like tissue outside the uterine cavity, typically within the pelvic peritoneum. Though traditionally explained by retrograde menstruation or stem cell misplacement, the persistence and recurrence of endometriotic lesions suggest a deeper regenerative misdirection. Chronic pelvic inflammation, hormonal dysregulation, and immune dysfunction alter the local tissue environment, promoting the differentiation of ectopic cells with endometrial characteristics. As these maladaptive phenotypes become dominant, they reinforce their own environment via estrogen production, angiogenesis, and inflammatory cytokines, exemplifying the threshold model of substitution.

## Barrett's Esophagus

In Barrett's esophagus, chronic gastroesophageal reflux leads to the replacement of normal squamous epithelium with columnar intestinal-type cells. This metaplastic transformation arises as a protective adaptation to acid exposure, yet results in increased cancer risk. Over time, the adaptive phenotype dominates the esophageal lining, disrupting regenerative fidelity and establishing a new, less functional baseline.

## Intestinal Metaplasia in Chronic Gastritis

Chronic infection with *Helicobacter pylori* or autoimmune gastritis induces inflammation and epithelial stress in the stomach lining. The gastric epithelium begins to express intestinal markers in an apparent attempt to survive persistent damage, illustrating the same pattern of loss of original phenotype and substitution by adaptive, yet inappropriate, cellular forms.

## Fibrotic Tissue Remodeling

In organs such as the liver, lungs, and pelvic peritoneum, chronic inflammation leads to excessive deposition of extracellular matrix and fibroblast activation. As fibrosis progresses, the normal parenchyma is gradually replaced by fibrotic tissue, not due to cell death alone but due to impaired regeneration. Fibrotic cells dominate the microenvironment, locking the tissue into a non-functional, maladaptive structure.

## Possible Extension to Early Neoplastic Transformation

In some pre-cancerous states, cells lose their differentiated identity and acquire stem-like, proliferative traits. This dedifferentiation may represent a final stage in the substitution process, where not only regenerative cues are lost, but control over growth and specialization is abandoned. Such transformations, particularly when linked to chronic inflammation and epigenetic instability, may be viewed as extensions of the same threshold model.

## Practical Implications

The Dominant Substitution Hypothesis not only offers a theoretical framework to reinterpret chronic degenerative diseases, but also suggests new practical strategies for intervention, diagnosis, and prevention. By focusing on the early phases of regenerative disruption, it may be possible to prevent maladaptive phenotypic dominance before the tipping point is reached.

## Early Identification of Regenerative Drift

Monitoring subtle shifts in cell phenotype, matrix composition, and inflammatory markers may allow clinicians to detect the onset of maladaptive regeneration. Advanced imaging, single-cell RNA sequencing, and tissue-specific epigenetic profiling could be leveraged to identify pre-substitution states in at-risk patients (Barker and Clevers 2010).

## Restoring the Original Regenerative Environment

Interventions aimed at re-establishing the structural and biochemical integrity of the regenerative niche may help maintain or recover original cellular identity. This includes modulation of the extracellular matrix, suppression of chronic inflammatory mediators, and hormonal balance restoration.

## Microbiota Modulation and Systemic Homeostasis

Given the systemic influence of gut microbiota on immune and metabolic signaling, strategies to correct dysbiosis—through diet, probiotics, prebiotics, or fecal microbiota transplant—may improve the cellular regenerative environment in distant tissues.

## Epigenetic Therapies and Differentiation Reprogramming

Pharmacological or nutrigenomic modulation of epigenetic regulators (e.g., DNA methyltransferase inhibitors, histone deacetylase inhibitors, methyl donors) could counteract maladaptive phenotypic fixation and promote re-differentiation toward the original cell type.

## Conceptual Shift in Chronic Disease Management

Rather than targeting end-stage symptoms or viewing disease as irreversible, this model supports a preventive and regenerative approach—focusing on preserving pattern fidelity, maintaining matrix integrity, and interrupting the cycle of substitution before functional collapse.

## Discussion

The Dominant Substitution Hypothesis provides a unifying framework to explain how chronic environmental disruption can reshape tissue identity through a progressive, proportion-based mechanism. This model shifts the focus from singular causative events (e.g., mutations, autoimmunity) to a system-level view where the interplay between inflammation, cellular adaptation, and loss of regenerative cues drives long-term dysfunction.

## Strengths of the Hypothesis

This model integrates findings from disparate fields—cell biology, immunology, epigenetics, and microbiome research—into a coherent explanation for a class of pathologies that share morphological and functional features. It also introduces a quantifiable concept: the threshold at which maladaptive phenotypes dominate. This may help explain why interventions are more effective at early stages and why tissue recovery becomes increasingly difficult once the original phenotype is marginalized.

## Limitations and Challenges

As with any theoretical model, empirical validation is essential. While supportive evidence exists in related domains, direct demonstration of substitution thresholds and proportion-driven degeneration requires longitudinal data, high-resolution tissue analysis, and well-controlled experimental models. Moreover, the hypothesis may not account for diseases with clearly monogenic origins or those triggered by acute, non-repetitive insults.

## Distinguishing Adaptation from Mutation

A critical clarification lies in differentiating epigenetically-driven phenotypic drift from irreversible mutational changes. The Dominant Substitution Hypothesis posits that many early changes are adaptive and reversible—given the right microenvironmental reset. This contrasts with models of disease that assume inevitable progression due to fixed genetic damage.

## Pathways for Experimental Testing

Future studies should aim to:

- Track regenerative phenotypes over time in models of chronic inflammation
- Quantify cell population ratios during disease onset and progression
- Manipulate the extracellular matrix, cytokine profiles, and microbiota to assess phenotypic reversibility
- Apply single-cell and spatial transcriptomics to detect early shifts in identity before morphological transformation

These experiments will be key to validating the existence of regenerative substitution thresholds and exploring therapeutic strategies aimed at pattern restoration rather than symptom suppression.

## Scope and Limitations of the Hypothesis

The Dominant Substitution Hypothesis is primarily applicable to tissues characterized by high cellular turnover and regenerative plasticity. These include epithelial surfaces (e.g., endometrium, intestinal mucosa, respiratory lining), mesothelial tissues (e.g., peritoneum), and organs with known capacity for cellular renewal (e.g., liver, skin). In such contexts, chronic environmental disruption interacts with ongoing regeneration, increasing the risk of maladaptive phenotypic replacement.

By contrast, the hypothesis does not apply to tissues with limited regenerative capacity, such as cardiac muscle, central nervous system neurons, or mature retinal tissue. In these organs, injury is more likely to result in cell death, scarring, or permanent loss of function, rather than adaptive phenotypic drift. Additionally, acute or rapidly resolved insults are unlikely to produce the slow threshold-based substitution described here.

Thus, this model should be understood as a framework for interpreting progressive tissue dysfunction in environments where chronic stress, attempted regeneration, and microenvironmental distortion co-exist over time. It does not aim to replace existing genetic or immunological theories of disease, but to complement them in specific scenarios of chronic cellular adaptation.

## Conclusions

The Dominant Substitution Hypothesis reinterprets chronic tissue degeneration as a progressive failure of regenerative fidelity, driven by environmental disruption and phenotypic drift. By shifting focus from isolated pathological events to systemic degradation of pattern guidance, this model provides a plausible mechanism for a wide range of conditions previously viewed as unrelated.

This framework challenges conventional interpretations of disease as purely mutational or autoimmune, offering a new lens grounded in regeneration dynamics and contextual cellular adaptation. If supported by experimental data, the hypothesis may pave the way for preventive strategies aimed at preserving cellular identity, reorganizing the extracellular matrix, restoring epigenetic stability, and modulating the microenvironment before irreversible substitution occurs.

Understanding degeneration as a result of the loss of original reference, rather than a random error or irreversible fate, reopens the discussion around reversibility, early diagnostics, and targeted regenerative interventions. The hypothesis lays conceptual groundwork for a regenerative medicine approach based not solely on cell replacement, but on the reactivation of lost guidance systems—the informational scaffolds that sustain tissue identity over time.

## Conflicts of interest

The authors have declared that no competing interests exist.

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