

The Role of Environmental Factors in Enhancing Immunological Responses among Celiac Disease Patients

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ABSTRACT

Celiac Disease (CD) is a widespread systemic autoimmune disorder triggered by dietary gluten in genetically predisposed individuals expressing HLA-DQ2/DQ8 genotypes. Because genetic susceptibility alone cannot account for the rising global incidence and diverse clinical phenotypes, current immunological research heavily focuses on external triggers. This study synthesizes peer-reviewed scientific literature to delineate the impact of environmental determinants on mucosal immunity and celiac disease pathogenesis, specifically addressing recent mechanistic advancements. Utilizing a rigorous methodological framework, this paper analyzes data from twenty-five landmark clinical and epidemiological studies across distinct primary domains: early infant feeding practices, gut microbiota dysbiosis, viral encounters, cellular T-cell dynamics, and industrial chemical pollutants. The synthesized evidence demonstrates that the breakdown of oral tolerance is a highly multifactorial process. Prolonged breastfeeding exerts protective mucosal effects by delivering secretory IgA (sIgA), whereas premature high-load gluten introduction overstimulates mucosal CD4⁺ T-cells. Concurrently, early-life antibiotic exposure induces severe gut dysbiosis—particularly depleting beneficial *Bifidobacterium* species—which compromises tight junctions, increases epithelial permeability, and triggers an evolutionary mismatch within historical microbial-cultural adaptation models. This mucosal vulnerability is often exacerbated by asymptomatic viral encounters, such as Reoviruses, which induce epigenetic modifications via mA RNA methylation, causing the immune system to misidentify dietary antigens as pathogens. At the cellular level, recent insights reveal strict tissue-mediated control, where enterocyte-derived IL-15 reprograms intraepithelial lymphocytes into aggressive cytotoxic cells, directly driving immunopathology. Moreover, modern industrial contaminants, including aluminum, act as powerful environmental adjuvants that disrupt epithelial integrity and lower the activation threshold for chronic autoimmunity. This synthesis concludes that celiac disease development relies on a cumulative interplay of genetic, epigenetic, and environmental pressures, emphasizing the critical necessity for early public screening and personalized preventative strategies in at-risk pediatric populations.

Keywords: Celiac disease; gluten; environmental factors; gut microbiota; breastfeeding

INTRODUCTION

BACKGROUND AND GLOBAL CONTEXT

Celiac Disease (CD) is a chronic enteropathy. It is an immune-mediated disorder. Dietary gluten triggers this condition. It develops in genetically susceptible individuals. Historically, it was considered a rare gastrointestinal disease. It mostly affected European pediatric populations. However, current estimates recognize it as a major public health concern. It affects about 1% of the worldwide population (Olsson et al., 2020). Diagnosed cases are rising absolutely. Additionally, incidence is surging in historical low-prevalence

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zones. These regions include the Middle East. They also include North Africa and parts of Asia (Cenit et al., 2021).

The underlying pathogenesis builds upon a profound immune response within the small intestinal lamina propria. This localized inflammation directly induces progressive villous atrophy and crypt hyperplasia. Ultimately, it leads to severe structural mucosal damage (Bouziat et al., 2017). This extensive enteropathy drives a highly diverse clinical spectrum. Patients frequently present with classical malabsorptive symptoms. These issues include chronic diarrhea and severe weight loss (Rubio-Tapia et al., 2013). Alternatively, individuals may manifest atypical extra-intestinal complications. These systemic manifestations include iron-deficiency anemia and chronic fatigue. Patients may also suffer from early-onset osteoporosis (Husby et al., 2020).

SOCIOECONOMIC, PSYCHOLOGICAL, AND CLINICAL BURDEN

The clinical management of celiac disease imposes a significant, lifelong burden on affected individuals. This challenge particularly impacts children and teenagers. Currently, the only validated therapeutic intervention is strict, lifelong adherence to a Gluten-Free Diet (GFD) (Namazy et al., 2022). This diet effectively restores mucosal health. Yet, a GFD presents immense practical challenges. Gluten-free products are expensive. They are also less accessible. These strict dietary restrictions cause social isolation. This isolation deeply affects younger patients during critical developmental periods. Recent data suggest a worrisome trend. Many patients experience persistent symptoms. They suffer from low-grade inflammation. This pathology occurs despite strict dietary compliance (Aronsson et al., 2019).

This vulnerability proves a major fact. Celiac disease is not a simple digestive reaction. Instead, it is a complex autoimmune disorder. This condition deeply impacts psychological well-being. It restricts dietary freedom. Ultimately, it reduces the overall quality of life for patients and their families (Aronsson et al., 2019).

THE INTRICATE INTERSECTION OF GENETICS AND ENVIRONMENT

From an immunological standpoint, celiac disease requires specific molecules. These are human leukocyte antigen (HLA) class II molecules. Specifically, they include HLA-DQ2 or HLA-DQ8 heterodimers. These molecules present gluten peptides to intestinal T-cells (Bouziat et al., 2017). These genotypes exist in nearly 30% to 40% of the global population. Yet, only a small fraction develops the disease. This represents approximately 2% to 3% of gene carriers (Antvorskov et al., 2018). This stark discrepancy proves a critical point. Genetic predisposition alone is an insufficient trigger. Over the last few decades, global incidence surged rapidly. This shift occurred too quickly for genetic mutations. Thus, it strongly points toward changing environmental determinants (Cenit et al., 2021).

Recent immunogenetic advances highlight intestinal barrier functions. Precise genetic frameworks govern these wall functions. These frameworks interact directly with global external stressors. Understanding specific HLA variations is essential. Non-HLA genetic risk factors are also crucial. Together, they map out mucosal destruction pathways. This explains why some individuals experience rapid damage while others remain asymptomatic (Aboulaghras et al., 2022). Therefore, external factors act as active modifiers. These factors include infant feeding regimens (Olsson et al., 2020) and industrial environmental pollutants (Namazy et al., 2022). They also include childhood antibiotic overuse (Aronsson et al., 2019) and mucosal viral infections (Bouziat et al., 2017). These elements disrupt early mucosal tolerance. They alter intestinal permeability. Ultimately, they drive the immune system into a destructive, chronic autoimmune cascade (Cenit et al., 2021).

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LITERATURE REVIEW

THE CHEMICAL STRUCTURE OF GLUTEN AND RESISTANCE TO PROTEOLYSIS

Gluten is a complex structural protein composite found in wheat, rye, and barley. This composite consists of two primary fractions, namely alcohol-soluble prolamins and alcohol-insoluble glutelins (Bouziat et al., 2017). The immunogenic potential of gluten is heavily concentrated within the toxic gliadin fraction. From a chemical perspective, this specific fraction is exceptionally rich in proline and glutamine amino acids (Antvorskov et al., 2018).

Due to this unique composition, the human gastrointestinal tract lacks specific post-proline endogenous proteolytic enzymes (Aronsson et al., 2019). Consequently, gluten undergoes incomplete digestion in the stomach and upper small intestine. This digestive failure leaves behind large, stable, and highly immunogenic peptide fragments in the lumen. The most notable of these triggers is the stable 33-mer peptide fragment. This structure successfully resists degradation by gastric, pancreatic, and brush-border membrane enzymes (Namazy et al., 2022). Ultimately, it reaches the intestinal mucosa entirely intact (Olsson et al., 2020). There, it initiates destructive inflammatory pathways within the tissue (Bouziat et al., 2017).

EARLY NUTRITIONAL PROGRAMMING: INFANT FEEDING AND BREASTFEEDING

Early dietary patterns can program the neonatal mucosal immune system. This biological hypothesis has gained substantial traction recently (Olsson et al., 2020). Historically, the timing of gluten introduction was thought to be the sole dietary variable influencing Celiac Disease pathogenesis. However, modern longitudinal cohort data have shifted the clinical focus toward a more nuanced view of early clinical markers (Cenit et al., 2021). These clinical findings demonstrated that early infant feeding practices directly modulate the baseline concentrations of anti-tissue transglutaminase (anti-tTG) autoantibodies (Aboulaghras et al., 2022). This modulation occurs well before clinical symptoms officially appear. This early nutritional programming is deeply tied to the protective role of breast milk. Prolonged breastfeeding serves as a multi-layered biological shield within the gut (Olsson et al., 2020).

Mother's milk introduces essential secretory IgA antibodies and anti-inflammatory cytokines. These protective elements effectively coat the infant's intestinal epithelium (Aronsson et al., 2019). Consequently, this coating chemically traps dietary antigens within the lumen. When infants are deprived of this maternal immunity, they face risks. Premature exposure to heavy gluten loads causes acute antigenic overload across the fragile mucosal barrier (Namazy et al., 2022).

This physiological stress hyperactivates naive CD4⁺ T-helper cells. This destructive pathway occurs before oral tolerance pathways can properly mature (Bouziat et al., 2017). Thus, it leaves the young child highly vulnerable to future autoimmune activation. Expanding upon this temporal framework, investigators shifted the lens to the prenatal period (Antvorskov et al., 2018). High maternal gluten intake during gestation modifies embryonic immune development within the womb. Ultimately, this prenatal exposure programs the fetal immune system to be more reactive after birth (Antvorskov et al., 2018).

MICROBIAL ARCHITECTURE AND CULTURAL ADAPTATION MODELS

Beyond direct nutrition, the microbial architecture of the human gut acts as a crucial intermediate layer (Olsson et al., 2020). This biological layer functions dynamically between dietary gluten and mucosal immunity (Bouziat et al., 2017). Beneficial gut bacteria produce vital short-chain fatty acids like butyrate. Specifically, *Bifidobacterium* species are major producers of these protective acids (Cenit et al., 2021). These short-chain fatty acids fuel enterocytes. They also maintain the structural integrity of tight junctions (Cenit et al., 2021). However, a loss of this microbial diversity undermines the entire barrier function (Aboulaghras et al., 2022). This modern loss is widely known as gut dysbiosis.

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When protective bacterial colonies decline, the tight junctions loosen. This damage creates a state of intestinal permeability or a leaky gut. This clinical state allows intact, immunogenic 33-mer gliadin peptides to pass easily into the lamina propria (Bouziat et al., 2017). Interestingly, this microbial vulnerability is frequently driven by broader cultural and industrial shifts. Modern human gluten tolerance required centuries of co-evolutionary microbial adaptation to agriculture (Cenit et al., 2021). Unfortunately, sudden modern shifts toward highly sterile living conditions severely disrupted this historical adaptation.

Industrialized food processing acts as a major external stressor in this process (Namazy et al., 2022). Furthermore, the chronic overuse of broad-spectrum antibiotics exacerbates this internal damage. This medical challenge is highly visible during early pediatric developmental windows (Aronsson et al., 2019). These environmental variables together create a profound evolutionary mismatch. Consequently, a damaged microbiome can no longer process modern dietary loads. Ultimately, this structural failure shifts the gut environment from healthy tolerance into an active chronic inflammation cascade (Cenit et al., 2021).

VIRAL INFECTIONS AND EPIGENETIC TRIGGERS

One critical breakthrough explains the sudden onset of celiac disease. This clinical onset occurs unexpectedly in genetically predisposed individuals (Cenit et al., 2021). Transient viral infections play a major role in this process (Namazy et al., 2022). Under normal conditions, the intestinal mucosa maintains a state of oral tolerance. Consequently, the healthy gut safely ignores harmless food proteins. However, a landmark study revealed a dangerous shift (Bouziat et al., 2017). Asymptomatic viral encounters can completely disrupt this established immunotolerance (Aronsson et al., 2019). Specifically, human Reoviruses are primary triggers of this mucosal disruption (Olsson et al., 2020).

During infection, the innate immune system launches an aggressive defense (Bouziat et al., 2017). This cellular defense releases high levels of pro-inflammatory cytokines. Type I Interferons are heavily produced during this viral response (Aboulaghras et al., 2022). Additionally, Interleukin-15 (IL-15) increases rapidly within the tissue. If gluten is ingested at this exact moment, problems arise (Bouziat et al., 2017). The hyper-activated immune system suffers a severe misclassification error. It mistakenly treats the harmless dietary protein as a dangerous viral pathogen (Namazy et al., 2022). This severe inflammatory confusion causes a permanent shift in immune memory.

Recent molecular research has uncovered the underlying epigenetic mechanisms. Dynamic RNA methylation heavily modulates these specific cellular pathways (Aboulaghras et al., 2022). This genetic process alters how the gut epithelium regulates antiviral responses. In celiac patients, these specific RNA modifications amplify viral-induced stress (Aboulaghras et al., 2022). This internal amplification prevents the tissue from returning to a calm state. Ultimately, it locks the mucosal immune system into a permanent loop. The child then remains trapped in a hyper-reactive autoimmune loop against gluten (Cenit et al., 2021).

ENVIRONMENTAL CONTAMINANTS AND ADVANCED IMMUNOLOGICAL ADJUVANTS

Diet, microbes, and viruses form the core of mucosal disruption (Olsson et al., 2020). However, modern industrial factors introduce an entirely different risk. These industrial variables add a dangerous chemical layer to celiac pathogenesis (Namazy et al., 2022). Exposure to industrial toxins and modern urban pollutants suppresses specific immune cells. Specifically, these contaminants prevent the normal development of regulatory T-cells (Cenit et al., 2021).

Consequently, this suppression prevents the immune system from learning how to properly regulate itself. This chemical threat can actually begin well before birth. Prenatal, in-utero exposure to environmental factors fundamentally alters early development (Antvorskov et al., 2018). This maternal exposure pre-programs the fetal immune system toward hypersensitivity. Thus, the biological damage occurs long before the child's first exposure to wheat (Antvorskov et al., 2018).

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Mechanistically, these modern toxins act as powerful immunological adjuvants. These chemical agents directly damage the fragile intestinal lining (Namazy et al., 2022). Patients with active celiac disease frequently exhibit elevated systemic immune responses. Common metals found in modern consumer products drive these environmental responses (Aboulaghras et al., 2022). Heavy metals can readily bind to human proteins within the body.

This binding causes structural changes that the immune system perceives as foreign threats. These chemical interactions induce severe cellular stress within the intestinal epithelium (Aboulaghras et al., 2022). This ongoing stress alters gene expression at a molecular level. Ultimately, these alterations lower the biological threshold required for gluten to trigger a full-scale autoimmune attack (Cenit et al., 2021).

MULTI-FACTORIAL RISK FACTORS FOR DISEASE DEVELOPMENT

The pathogenesis of celiac disease is fundamentally non-linear. It represents a complex biological paradigm. Genetic susceptibility alone is clearly insufficient to trigger clinical manifestation (Aboulaghras et al., 2022). Disease development depends on a dynamic multi-factorial interplay. Genetic predisposition converges with cumulative environmental pressures to breach immune tolerance (Cenit et al., 2021). Early nutritional programming also plays a key role in this breakdown. Dietary gluten serves as the mandatory primary trigger. However, secondary environmental orchestrators heavily dictate the actual clinical onset. These factors include the volume and timing of early childhood gluten intake (Olsson et al., 2020). Additionally, maternal dietary factors during gestation influence this risk (Antvorskov et al., 2018).

Modern epidemiological data highlights several overlapping risks. Infant feeding strategies and early antibiotic-induced gut dysbiosis act synergistically (Aronsson et al., 2019). Furthermore, transient viral encounters and industrial chemical contaminants compound this damage (Bouziat et al., 2017). These various elements do not act in isolation (Cenit et al., 2021). Together, they progressively compromise intestinal tight junctions. This structural failure lowers the activation threshold for mucosal immune cells (Aboulaghras et al., 2022).

Consequently, modern industrialized lifestyles and chemical exposure create a profound evolutionary mismatch (Namazy et al., 2022). This ongoing exposure shifts a controlled dietary intake into a destructive autoimmune cascade. Therefore, understanding celiac disease requires a holistic model. This medical model must evaluate how diverse risk factors collectively disrupt host immunity over time (Namazy et al., 2022).

IMMUNOLOGICAL PATHWAYS OF CELIAC DISEASE

The pathogenesis of Celiac Disease represents a classic biological intersection. It marks a critical meeting point between innate and adaptive immunity. Large, indigestible gliadin peptides easily cross the weakened epithelial barrier (Aboulaghras et al., 2022). These foreign proteins then directly enter the lamina propria. Once inside, they initiate a highly coordinated and destructive immunological cascade (Bouziat et al., 2017). This pathological process can be broken down into four critical physiological phases:

ENZYMATIC MODIFICATION VIA TISSUE TRANSGLUTAMINASE (TTG)

Gliadin fragments easily gain access to the lamina propria of the small intestine. The stable 33-mer peptide is a primary example of these fragments. Once inside, they promptly encounter the endogenous enzyme tissue transglutaminase 2 (tTG-2). Under normal physiological conditions, tTG-2 is involved in tissue repair and cross-linking (Cenit et al., 2021). However, a dangerous shift occurs in celiac disease. The enzyme targets the proline-and-glutamine-rich gliadin fragments for enzymatic modification (Aboulaghras et al., 2022). This specific chemical process is known as deamidation.

During deamidation, tTG-2 converts specific neutral glutamine residues within the peptide sequence. These residues transform into negatively charged glutamic acid (Aboulaghras et al., 2022). This biochemical transformation is the critical turning point in the disease's development. The introduction of these negative charges drastically increases the chemical affinity of the gliadin peptides. Consequently, they fit perfectly

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into the positively charged antigen-binding grooves of human immune cells (Cenit et al., 2021). These specific grooves are present on the HLA class II molecules.

ADAPTIVE IMMUNE ACTIVATION AND T-CELL DYNAMICS

Enzymatic modification changes the gliadin peptides. Following this process, professional Antigen-Presenting Cells (APCs) immediately recognize and engulf these negatively charged deamidated fragments. These specific APCs express HLA-DQ2 or HLA-DQ8 genotypes (Aboulaghras et al., 2022). The APCs then present these immunogenic complexes to naïve CD4⁺ T-helper cells within the lamina propria. This specific presentation initiates a rapid clonal expansion of gluten-reactive CD4⁺ T-cells (Cenit et al., 2021). Once activated, these T-helper cells act as the primary drivers of chronic immunopathology. They effectively convert a localized dietary reaction into a systemic autoimmune cascade (Aboulaghras et al., 2022).

These activated CD4⁺ T-cells proliferate rapidly. This cellular proliferation secretes a massive wave of pro-inflammatory Th1-associated cytokines. Primarily, this wave consists of Interferon-gamma (IFN-gamma) and Tumor Necrosis Factor-alpha (TNF-alpha) (Aboulaghras et al., 2022). These powerful cytokines act directly on the surrounding environment. They dismantle epithelial tight junctions from the basolateral side. Consequently, this structural damage drastically accelerates mucosal permeability. Furthermore, this intense cytokine storm drives resident B-lymphocytes to differentiate into specialized plasma cells. This differentiation initiates the synthesis of highly specific anti-tissue transglutaminase (anti-tTG) and anti-deamidated gliadin peptide (anti-DGP) autoantibodies (Namazy et al., 2022). Crucially, this adaptive T-cell response does not act in isolation. Its destructive and autoimmune potential is strictly gated, amplified, and maintained by stress signals. These signals come directly from the surrounding intestinal epithelial tissue environment (Cenit et al., 2021).

INNATE TISSUE CONTROL, INFLAMMATORY CASCADE, AND AUTOIMMUNITY

Simultaneously, a profound innate immune cascade occurs within the epithelial layer. This process represents a classic example of tissue-driven autoimmunity. Stressed enterocytes respond to specific environmental insults. Consequently, these cells overproduce Interleukin-15 (IL-15). This particular cytokine acts as a master regulator of tissue pathology (Bouziat et al., 2017). This localized overproduction of IL-15 directly reprograms Intraepithelial Lymphocytes (IELs). This chemical reprogramming transforms normal cytotoxic CD8⁺ T-cells and Natural Killer (NK) cells. As a result, they become aggressive, antigen-independent killers. These modified cells directly target the host's own tissue (Aboulaghras et al., 2022).

Under this tissue-mediated signaling, IELs rapidly upregulate natural killer receptors. Specifically, they upregulate NKG2D receptors. These receptors bind tightly to stress-induced ligands (MICA/MICB) on the cell surface (Aboulaghras et al., 2022). These specific ligands are expressed on the damaged surface of the enterocytes. This molecular interaction triggers a massive, direct cytotoxic attack on the epithelial cells. This uninhibited destruction drives the cells into widespread, accelerated apoptosis (Bouziat et al., 2017). The cumulative result of this destruction is the classic histopathological hallmark of celiac disease. Patients exhibit severe villous atrophy and crypt hyperplasia. This destructive cascade proves that celiac disease is not just a localized food allergy. Instead, it is a true tissue-mediated autoimmune destruction. The intestinal epithelium actively dictates its own pathological fate (Aboulaghras et al., 2022).

EPIGENETIC MODULATIONS AND CELLULAR RECEPTORS (M6A AND CD247)

Modern molecular biology has unveiled complex epigenetic layers. These mechanisms go far beyond classical T-cell activation. These biological layers control how the host immune system responds (Cenit et al., 2021). Specifically, dynamic RNA modifications like m6A methylation play a fundamental role within the gut epithelium of celiac patients. These molecular changes heavily modulate crucial antiviral pathways (Aboulaghras et al., 2022). When a viral trigger is present, these RNA modifications can quickly amplify the inflammatory state. Consequently, this heightened inflammation prevents the intestinal tissue from returning to normal homeostasis (Olsson et al., 2020).

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In a strict molecular context, structural variations in the CD247 gene alter immune responses. This gene encodes a vital component of the essential T-cell receptor complex. These genetic variations directly interact with external chemical and viral exposures (Cenit et al., 2021). This ongoing interaction progressively compromises cellular defense mechanisms. Ultimately, these molecular alterations lower the biological threshold required to trigger chronic autoimmunity (Namazy et al., 2022).

METHODOLOGY

To systematically investigate the multi-layered impact of environmental determinants on mucosal immunity in celiac disease, a rigorous review and critical synthesis of landmark clinical trials, longitudinal birth cohorts, and mechanistic profiles were conducted. A total of twenty-five peer-reviewed papers were critically selected to construct a comprehensive operational overview. To enhance theoretical cohesion, the essential parameters evaluated across these twenty-five sources are systematically synthesized in Table4.1 below, sequentially structured to mirror the clinical timeline—advancing from early gestational programming and environmental/microbial triggers to downstream cellular cytotoxicity and global geographic disparities:"

Table 4.1. Detailed methodological breakdown of celiac disease variables

Author & Year	Environmental / Biological Trigger	Cellular Immune Mechanism	Histopathological / Lab Findings	Critical / Geographic Gaps
Antvorskov et al. (2018)	Maternal Gluten Load & Gestational Diet	Intrauterine immune priming of the fetal immune system	Alteration in baseline pediatric primary biomarkers at birth	Scarcity of multi-generational longitudinal tracking of mothers
Unalp-Arida et al. (2014)	Early Nutritional Programming	Activation of early immune response against gluten antigens	Distinct dynamics in anti-tTG antibody concentrations	Gaps in tracking pediatric baseline biomarkers pre-clinically
Hill et al. (2022)	Continuous Gluten Exposure	Activation of tissue transglutaminase antibodies	Elevated serum levels of anti-tTG antibodies	Diagnostic gaps in silent or asymptomatic cohorts
Karvonen et al. (2023)	Gluten-Free Diet (GFD) Elimination	Gradual resolution of mucosal immune inflammation	Tracking of long-term mucosal healing timelines	High variability in tissue recovery rates among patients
Desrosiers et al. (2023)	Primary Cellular Triggers	Initiation of autoimmune attack against enterocytes	Structural transformations leading to early mucosal atrophy	Lack of reference baseline structural data for early-stage atrophy
Turner & Bernstein (2022)	Chronic Inflammation	Increased paracellular permeability via zonulin pathways	Degradation and compromise of tight junction barrier proteins	Complete paracellular permeability mechanisms remain unclear

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Bouziat et al. (2017)	Transient Viral Infection (Reovirus)	Induction of a mucosal 'misclassification error'	Breakdown of oral tolerance toward dietary gliadin antigens	Failure to identify all specific viruses causing this error
Sebastian-delaCruz et al. (2025)	Cellular & Viral Induced Stress	Activation of dynamic m6A RNA epigenetic methylation loops	Acceleration of mucosal autoimmune inflammatory cascades	Lack of global validation of this specific epigenetic mechanism
Namazzy et al. (2022)	Sterile Living Environments	Failure of regulatory T-cell (Treg) maturation	Loss of systemic immune balance and dysregulation	Need for long-term comprehensive reviews of the hygiene hypothesis
Catassi et al. (2024)	Early Childhood Antibiotic Overuse	Induction of profound gut microbiota imbalance (Dysbiosis)	Fundamental compromise of intestinal epithelial barrier integrity	Lack of direct dose-response studies linking antibiotics to celiac onset
Bektaş & Ulusoy (2025)	Modernization & Industrialized Lifestyle	Induction of an evolutionary mismatch	Increased incidence rates of intestinal autoimmune activation	Insufficient study on the impact of modern environments in specific cohorts
Abadie & Jabri (2014)	Epithelial Chronic Stress	Massive overproduction of Interleukin-15 (IL-15) as a survival factor	Reprogramming and activation of intraepithelial lymphocytes (IELs)	Historical absence of targeted therapeutic interventions for this pathway
Asri et al. (2021)	Mucosal Hyper-responsiveness	Perpetuation of an autoimmune loop expanding beyond gluten	Extensive, widespread, and systemic destruction of gut architecture	Insufficiency of strict dietary gluten exclusion alone
Mearin & Jabri (2024)	Chronic IL-15 Signaling	Reprogramming of IELs into aggressive natural killer-like cells	Induction of widespread lymphocyte reprogramming pathways	Clinical lack of standardized, specific IL-15 inhibitors
Rahmani et al. (2025)	Stress-Induced Cellular Ligands	Selective upregulation of cytotoxic NKG2D receptors	Triggering of extensive enterocyte apoptosis	Shortage of interventions to halt cell death before structural atrophy

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Barada et al. (2014)	Regional Epidemiological Factors	Immune activation linked to local genetic backgrounds	Documentation of distinct variations in prevalence and phenotypes	Critical geographic gap across the Middle East and North Africa
Rostami et al. (2018)	Diagnostic Infrastructure Limits	Delayed screening activation due to absence of clinical guidelines	Representation of a diagnostic paradox (high HLA frequency vs. low documentation)	Severe lack of standardized serological screening protocols in developing regions
Olsson et al. (2020)	Western Screening Strategies	Early clinical immune surveillance and prompt screening	Ideal documentation of pediatric clinical timelines	Absence of these structured screening models in non-Western areas
Long et al. (2021)	Socioeconomic Status & Urbanization	Material and environmental impacts on disease manifestation	Underreported official data and idealized Western screening templates	Inaccurate global prevalence models due to socioeconomic disparities
Karthikeyan et al. (2024)	Variable Serological Protocols	Inefficient antibody monitoring due to unstandardized testing	High disparity in laboratory readings across non-Western cohorts	Absence of universally standardized, resource-specific serological tools
Catassi & Fasano (2024)	Global Environmental Stressors	Global shifts in patterns of local mucosal immune tolerance	Surging global incidence rates and shifting disease paradigms	Current predictive models fail to include diverse global phenotypes
Aronsson et al. (2019)	Timing of Pediatric Gluten Introduction	Interplay between gut microbial environment and timing of antigen exposure	Large-scale screening data within the pediatric TEDDY clinical trial	Severe scarcity of multi-omic longitudinal studies tracking non-Western cohorts
García-Santisteban et al. (2020)	Microbial Programming	Microbial composition impact on epithelial tight junction stability	Alterations in intestinal epithelial barrier integrity in children	Lack of preventative strategies based on early-life microbial programming
Visvanathan et al.	Epigenetics	Complex multi-	Deep molecular	Profound lack of

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(2021)	Regulations	omic interaction in autoimmune enteropathies	and biological insights into intestinal inflammation	diverse, ethnically varied epithelial cell lines in lab models
Cenit et al. (2021)	Industrialization Mismatch	Alteration of immune tolerance thresholds via environmental chemical exposure	Epigenetic alterations and variations in local gene expression	Scarcity of direct data linking specific industrial pollutants to shifts in tolerance

DISCUSSION

COMPARATIVE ANALYSIS OF DIETARY TRIGGERS AND MUCOSAL MATURATION

Early-life nutritional exposures actively influence the transition from genetic susceptibility to clinically manifest celiac disease. Infant feeding practices contribute significantly to shaping mucosal immunity during critical periods of immune maturation (Antvorskov et al., 2018). Breast milk, in particular, delivers vital bioactive compounds and secretory immunoglobulin A (sIgA) to the infant gut. These components limit the interaction between dietary antigens and the intestinal epithelium, thereby supporting oral tolerance and preventing excessive immune activation (Olsson et al., 2020).

The timing and quantity of gluten introduction also serve as critical determinants of subsequent disease risk. High gluten intake during early windows of incomplete immune development can accelerate antigenic stimulation and provoke inflammatory pathways. Specifically, excessive early exposure facilitates the activation of gluten-reactive CD4+ T lymphocytes (Olsson et al., 2020). This activation often occurs before the mucosal barrier establishes robust regulatory mechanisms, increasing the likelihood of tissue damage in genetically predisposed infants (Lionetti et al., 2014).

Importantly, these dietary influences may initiate during the prenatal period through maternal nutrition. Maternal dietary habits during pregnancy function as early modulators of fetal immune programming. High maternal gluten consumption can elevate fetal exposure to dietary antigens or circulating inflammatory mediators, altering immune responsiveness after birth (Antvorskov et al., 2018). Longitudinal cohort data confirm this link, showing direct associations between early nutritional environments and elevated levels of celiac-related serological markers, including anti-tissue transglutaminase antibodies (anti-tTG) (Aronsson et al., 2019).

However, clinical findings regarding the preventive efficacy of breastfeeding and managed gluten introduction remain inconsistent. While certain prospective studies support a protective effect, several major clinical trials report no significant reduction in disease incidence (Lionetti et al., 2014; Chmielewska et al., 2015). These discrepancies stem from variations in study design, cohort genetics, and early-life microbiome composition (Aronsson et al., 2015). Consequently, prenatal and postnatal nutritional factors do not act in isolation; instead, they interact dynamically with genetic risks to govern long-term immune tolerance to gluten.

MICROBIAL DISRUPTION, INTESTINAL PERMEABILITY, AND MODERN LIFESTYLE FACTORS

The gut microbiome plays a foundational role in maintaining intestinal homeostasis and modulating immune pathways linked to celiac disease (Cenit et al., 2021). Under physiological conditions, a diverse microbial

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ecosystem preserves epithelial barrier integrity and promotes oral tolerance (Olsson et al., 2020). Beneficial bacterial genera, particularly *Bifidobacterium* species, ferment dietary fibers to produce short-chain fatty acids (SCFAs). These metabolites strengthen tight junction stability and drive anti-inflammatory signaling pathways, effectively preventing the systemic translocation of luminal antigens (Cenit et al., 2021).

Conversely, microbial dysbiosis compromises epithelial barrier structures and alters host immunoregulation. Longitudinal data indicate that a marked reduction in these beneficial bacterial populations often precedes or accompanies the onset of celiac disease clinical markers (Aronsson et al., 2019). This taxonomic shift impairs tight junction complexes, facilitating the paracellular passage of immunogenic gluten peptides into the intestinal mucosa. Once inside, these peptides interact directly with antigen-presenting cells to trigger localized inflammatory cascades (Olsson et al., 2020).

Early-life environmental insults, particularly the overuse of broad-spectrum antibiotics, heavily drive these microbial disruptions. Antibiotic exposure during critical developmental windows depletes microbial diversity and arrests the maturation of a stable intestinal ecosystem (Aronsson et al., 2019). This resulting imbalance suppresses regulatory T-cell (Treg) differentiation, thereby undermining the baseline mechanisms required to maintain immune tolerance to dietary proteins (Olsson et al., 2020).

Modern industrialized lifestyles further accelerate mucosal stress through ultra-processed diets rich in food additives and emulsifiers. These dietary components alter the mucosal layer, inducing low-grade intestinal inflammation and increasing epithelial permeability (Cenit et al., 2021). While these environmental shifts closely parallel the rising global incidence of celiac disease, establishing definitive causal relationships remains a challenge. Consequently, future longitudinal and mechanistic evaluations within established cohorts are essential to clarify whether these microbial alterations represent primary drivers of disease pathogenesis or secondary consequences of ongoing mucosal inflammation.

VIRAL OVERLAP, ORAL TOLERANCE BREAKDOWN, AND THE HYGIENE HYPOTHESIS

Specific viral infections actively disrupt the immunological mechanisms governing oral tolerance to dietary proteins. Transient viral exposures during critical windows of mucosal maturation can alter host responses to dietary antigens, thereby accelerating the transition toward pathogenic auto-reactivity (Cenit et al., 2021). Pathogens such as reoviruses and enteroviruses induce localized pro-inflammatory signaling that effectively blocks the default tolerogenic pathways in the gut. This viral-mediated interference prompts the immune system to misidentify dietary gliadin as a hostile threat rather than a harmless nutrient (Olsson et al., 2020).

This pathogenic viral overlap aligns closely with the principles of the hygiene hypothesis. In highly industrialized environments, diminished exposure to diverse microbial stimuli alters early immunoregulatory pathways (Cenit et al., 2021). This lack of microbial priming impairs the functional development and suppressive capacity of regulatory T cells (Tregs), which are essential for maintaining long-term mucosal tolerance. Consequently, when a virus triggers an inflammatory response in an under-primed immune system, the baseline mechanisms required to suppress anti-gluten responses fail to operate efficiently (Olsson et al., 2020).

.At the molecular level, viral-induced epithelial stress triggers complex intracellular changes, including post-transcriptional modifications within the intestinal mucosa. Epigenetic and m^6A RNA modifications participate directly in regulating these localized inflammatory pathways, serving as molecular rheostats that dictate the intensity of the mucosal immune response (Aboulaghras et al., 2022). These subtle molecular alterations further compromise the epithelial barrier, facilitating the paracellular translocation of immunogenic gluten peptides into the lamina propria where they encounter antigen-presenting cells (Aronsson et al., 2019). Viral exposures, however, do not prompt clinical pathogenesis in isolation; instead, they function within a synergistic network of environmental insults. Concomitant early-life factors, such as broad-spectrum antibiotic use, induce a state of dysbiosis that severely impairs baseline intestinal barrier function (Aronsson et al., 2019). Therefore, the convergence of microbial dysbiosis, viral stress, and genetic susceptibility collectively drives the breakdown of oral tolerance. This multi-hit framework suggests that

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viral infections act as kinetic triggers rather than independent causes, accelerating celiac disease onset only when the mucosal environment is already compromised.

GEOGRAPHIC VARIATIONS AND CRITICAL RESEARCH GAPS

The global epidemiology of celiac disease exhibits profound geographic heterogeneity, driven by a complex interplay between population genetics and shifting environmental exposures. Marked disparities in prevalence rates and clinical presentation exist between Western nations and developing regions, including the Middle East and North Africa (Cenit et al., 2021). While higher diagnosis rates in Western cohorts reflect widespread clinical awareness and robust healthcare infrastructure, underdiagnosis remains a pervasive challenge in developing nations (Cenit et al., 2021). This diagnostic discrepancy becomes especially critical given that several non-Western populations possess a remarkably high prevalence of HLA susceptibility alleles. Despite this strong genetic predisposition, reported clinical disease rates remain paradoxically low, a mismatch indicating that limited diagnostic infrastructure, restricted access to serological testing, and the lack of unified screening strategies severely underestimate the true global disease burden (Olsson et al., 2020).

In tandem, rapid urbanization and modern dietary shifts alter host immunity and change how the disease presents clinically across different regions (Olsson et al., 2020). These geographic imbalances highlight several critical research gaps that obstruct a comprehensive understanding of celiac disease pathogenesis. Available data derive predominantly from homogeneous Western cohorts, while longitudinal initiatives tracking ethnically diverse populations remain scarce. To resolve these limitations, prospective birth cohorts must follow high-risk individuals from gestation through early childhood. Such frameworks are vital to map the precise mechanistic interactions among maternal dietary factors, early-life microbiome assembly, and mucosal immune maturation (Aronsson et al., 2019; Olsson et al., 2020).

Moreover, current experimental and laboratory models possess inherent mechanistic limitations. While established in vitro designs offer valuable insights into cellular stress, they often fail to capture the broader genetic, epigenetic, and ethnic diversity seen globally. Expanding the application of diverse cellular systems is therefore essential to clarify the post-transcriptional and immunological pathways driving mucosal damage across different populations (Aboulaghras et al., 2022).

Ultimately, celiac disease cannot be evaluated through a single, static epidemiological framework. Geographic, genetic, and socioeconomic realities collectively shape both risk profiles and diagnostic outcomes. Addressing these critical knowledge gaps through internationally coordinated, ethnically diverse cohorts is not merely a methodological necessity; it is the definitive path to refining global preventive strategies and ensuring equitable diagnostic accuracy worldwide.

CONCLUSION & RECOMMENDATIONS

CONCLUSION

This review highlights the multifactorial nature of celiac disease and emphasizes the complex interactions between genetic susceptibility and environmental influences in disease development. Although the presence of susceptible HLA genotypes is a prerequisite for disease occurrence, genetic factors alone cannot fully explain the increasing global prevalence of celiac disease.

Current evidence suggests that a variety of environmental exposures, including early-life nutritional factors, alterations in the gut microbiome, antibiotic use, viral infections, and environmental pollutants, may contribute to the disruption of immune tolerance to gluten in genetically predisposed individuals. These factors appear to influence intestinal barrier integrity, microbial composition, and immune regulation, thereby creating conditions that may facilitate the initiation and progression of disease-related immune responses.

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emerging research indicates that environmental factors may exert their effects through complex molecular and epigenetic mechanisms that remain incompletely understood. The interaction between these pathways and host immune responses represents an important area for future investigation.

Overall, celiac disease should be viewed as the result of dynamic interactions among genetic, environmental, microbial, and immunological factors rather than as a purely genetic disorder. A deeper understanding of these interactions may contribute to improved risk assessment, earlier disease detection, and the development of more effective preventive and therapeutic strategies in the future.

FUTURE RECOMMENDATIONS

Translating the current mechanistic evidence into actionable clinical and public health outcomes requires targeted, strategic interventions across research, policy, and clinical practice: **Optimizing Early-Life Nutritional Frameworks:** Public health policies must actively prioritize and support prolonged breastfeeding alongside evidence-based infant feeding guidelines. These early dietary frameworks are crucial to guide physiological immune maturation and secure oral tolerance networks during critical developmental windows.

Enforcing Pediatric Antimicrobial Stewardship: Healthcare providers must rigorously enforce the rational prescribing of broad-spectrum antibiotics during infancy. Implementing strict antimicrobial stewardship programs within pediatric care is essential to prevent the premature depletion of protective microbial taxa, thereby preserving gut homeostasis.

Expanding Longitudinal and Diverse Cohorts: Future epidemiological funding should prioritize large-scale, prospective birth cohorts that follow high-risk individuals from gestation through childhood. Crucially, these initiatives must expand into historically underrepresented regions, specifically the Middle East and North Africa, where high genetic susceptibility contrasts sharply with sparse systematic screening data.

Integrating Multi-Omic and Epigenetic Frameworks: Advanced research designs should adopt multidisciplinary pipelines that converge immunology, microbiology, and post-transcriptional biochemistry. Investigating localized mucosal alterations—particularly m6A RNA modifications—will be instrumental in discovering early, non-invasive biomarkers and identifying novel non-dietary therapeutic targets.

Upgrading Diagnostic Infrastructure and Global Awareness: Reducing the global underdiagnosis burden requires a unified effort to strengthen clinical infrastructure, expand access to standardized serological screening, and raise public health awareness. Enhancing these diagnostic capabilities is the definitive step toward achieving early clinical detection and establishing personalized preventive monitoring for genetically vulnerable individuals.

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