

# **Trauma-Induced Epigenetic Modifications and Gene Expression**

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## Abstract

Traumatic experiences—physical, psychological, or sociopolitical—can leave durable “molecular scars” that tune the activity of hundreds to thousands of genes without mutating DNA. These epigenetic changes are now linked to altered stress-hormone signalling, immune dysregulation, neuroplasticity, accelerated biological ageing and, in some cases, transmission of vulnerability to the next generation. Recent high-resolution epigenome-wide association studies (EWAS) and multi-omic approaches confirm that trauma reshapes DNA-methylation landscapes, histone marks, chromatin accessibility, small-RNA profiles, and m<sup>6</sup>A RNA methylation in a time-, tissue- and sex-specific manner. Therapeutically, reversible epigenetic marks offer an attractive target for psychotherapy-assisted pharmaco-epigenetic, lifestyle, and nutraceutical interventions.

## Introduction

### Epigenetic mechanisms in brief

Mechanism	Functional effect	Trauma-responsive examples
DNA methylation (5-mC)	Silences or fine-tunes gene transcription	NR3C1, FKBP5, BDNF, AHRR, DUSP22
Histone post-translational modifications (acetylation, methylation, phosphorylation)	Alter chromatin compaction and recruitment of transcription machinery	H3K27ac/H3K9ac loss in prefrontal cortex after chronic stress; restoration with HDAC inhibitors
Chromatin remodelling	Nucleosome repositioning controls promoter/enhancer accessibility	Large DMR blocks in inflammatory-coagulation genes after severe injury
Non-coding RNAs (miRNA, lncRNA, circRNA)	Post-transcriptional repression, sponging, chromatin targeting	miR-16, miR-125a, miR-124, lncRNA NEAT1 dysregulated in PTSD
Epitranscriptomic marks (m <sup>6</sup> A)	Controls mRNA stability and translation	FTO-driven m <sup>6</sup> A demethylation up-regulates BDNF-TrkB pathway in memory reconsolidation after stress

## Background or Context

### Trauma typology and critical windows

- **Prenatal & pre-conception stress** – Maternal PTSD symptoms or war exposure alter placental NR3C1 and LEP methylation; sperm DNAm at immune genes predicts offspring neurodevelopment.
- **Early-life adversity (0-5 y)** – ACE scores correlate with genome-wide methylation shifts decades later, especially in stress-immune loci.
- **Adolescence/adulthood trauma** – Acute physical injury triggers >10 000 differentially methylated CpGs within 60 min that partly persist one month later.
- **Chronic or cumulative trauma** – First-responder meta-analysis shows dose-dependent hypomethylation at HPA-axis genes across three trauma-related phenotypes.

## Main Argument or Methods

### Intergenerational and transgenerational findings

- **Human evidence** – 2025 Nature Scientific Reports analysis of 126 Syrian families documents 21 stress-related CpG changes in mothers and 14 persisting to grandchildren, including accelerated epigenetic ageing clocks.
- **Animal models** – Paternal predator-odor stress alters sperm miRNA cargo and primes F1 hippocampal fear circuits; stress-induced lncRNA changes propagate chronic-pain risk.

### Health implications

Domain	Epigenetic link	Clinical outcome
Mental health	NR3C1 / FKBP5 methylation, miR-124 dysregulation	PTSD, depression, anxiety, suicidality
Cardiometabolic	Hypo-methylation at inflammatory and adipokine promoters	Hypertension, metabolic syndrome, atherosclerosis
Immune & pain	DMRs in PAX8, DUSP22; cytokine miRNA profiles	Auto-immune disorders, chronic pain syndromes
Neurocognitive	BDNF epigenetic repression, m <sup>6</sup> A imbalance	Memory deficits, executive dysfunction
Oncogenesis	Stress-induced loss of tumor-suppressor methylation	Cancer progression potential (mechanistic models)

## Analysis or Case Studies

### Molecular evidence linking trauma to gene expression

Gene / pathway	Epigenetic alteration	Functional consequence	Evidence
<b>NR3C1 (glucocorticoid receptor)</b>	Promoter hyper-methylation	Blunted cortisol feedback, prolonged HPA axis activation	PTSD psychotherapy responders show demethylation & symptom improvement
<b>FKBP5 (HPA co-chaperone)</b>	Trauma-related intronic CpG hypo-methylation inherited in offspring	Enhanced stress hormone sensitivity & risk for depression/PTSD	Holocaust and war-refugee cohorts; Syrian 3-generation study
<b>BDNF</b>	Context-specific promoter methylation; m <sup>6</sup> A-mediated up-regulation	Impaired synaptic plasticity, learning, mood	Rodent predator-stress model; human blood EWAS
<b>Immune / coagulation genes (e.g., AEBP2, PPP1CA)</b>	Large DMR blocks after polytrauma	Persistent inflammation, infection risk	Prospective trauma cohort
<b>miRNAs (e.g., miR-16, miR-125a)</b>	Up- or down-regulation	Targets serotonergic, immune and apoptosis genes	Clinical miRNA panels in veterans

## Conclusion

### Reversibility and therapeutic opportunities

#### 1. Psychotherapy & mind-body programmes

- Evidence-based CBT, EMDR and mindfulness-based stress reduction normalize FKBP5 methylation and diurnal cortisol in responders.

#### 2. Pharmaco-epigenetic agents

- **HDAC inhibitors (e.g., valproic acid, SAHA)** promote fear-extinction gene expression; sex-specific efficacy shown in rat PTSD models.
- **DNMT inhibitors (e.g., RG-108)** transiently disrupt traumatic memory reconsolidation in mice; safety in humans remains under study.

#### 3. Lifestyle & nutraceuticals

- B-vitamin complexes mitigate hippocampal mtDNA methylation and PTSD-like behaviours in rodents.
- Physical activity, anti-inflammatory diet and social support associated with beneficial epigenetic ageing trajectories after adversity.

## Methodological challenges

- **Tissue specificity** – Blood profiles may not mirror brain circuits driving psychopathology.
- **Causal inference** – Reverse causation (disease changing behaviour) and gene–environment correlation complicate analyses.
- **Heterogeneity of trauma and timing** – Need for harmonised exposure metrics and longitudinal designs.
- **Ethics & privacy** – Epigenetic data can reveal personal history of abuse; governance frameworks must protect participants.

## Future directions (2025-2030)

1. Multi-omic single-cell atlases tracking chromatin, methylome and transcriptome after trauma.
2. CRISPR-dCas-based locus-specific epi-editing to reverse maladaptive marks without global off-target effects.
3. Large-scale randomised trials combining psychotherapy with targeted HDAC/ DNMT modulators.
4. Policy translation: integrating epigenetic screening with psychosocial services for conflict and disaster zones.

## References

1. *Systematic review of gene expression and epigenetics in first-responders* (2025). <https://www.sciencedirect.com/science/article/>
2. *Nature npj Genomic Medicine: Severe traumatic injury methylome study* (2024). <https://www.nature.com/articles/>
3. *NR3C1 methylation and PTSD therapy outcomes* (2023). <https://www.nature.com/articles/>
4. *Syrian refugee three-generation DNA-methylation study* (2025). <https://www.nature.com/articles/>
5. *Valproic-acid HDAC-inhibitor PTSD model* (2024). <https://pubmed.ncbi.nlm.nih.gov/>
6. *miRNA biomarkers in PTSD review* (2024). <https://pubmed.ncbi.nlm.nih.gov/>
7. *Framework of epigenetic pathways to resilience* (2025). <https://pubmed.ncbi.nlm.nih.gov/>
8. *Exposure to war and methylation mediation of PTSD symptoms* (2023). <https://www.frontiersin.org/journals/epidemiology/articles/>