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Research Paper

Decoding Short- and Long-Term Cellular Adaptations to Cr(VI) Exposure Through High-Throughput Transcriptomics

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Abstract

Hexavalent chromium (Cr(VI)) is a well-established environmental and occupational carcinogen, but its time-dependent molecular effects remain poorly characterized. This study aims to elucidate the transcriptional responses triggered by acute versus chronic Cr(VI) exposure through an integrated analysis of two publicly available transcriptomic datasets: GSE16349 (short-term exposure, 16 hours) and GSE24025 (long-term exposure, 4 weeks). We identified 250 differentially expressed genes across both exposure models. MetaCore pathway enrichment analysis revealed shared activation of apoptosis, survival signaling, DNA damage response and repair, and cell cycle progression. Notably, short-term exposure primarily activated acute stress responses, whereas long-term exposure induced reprograms transcription toward fibrosis, EMT, and oncogenic signaling. Protein-protein interaction (PPI) network analysis identified potential key hub genes, with potential as biomarkers for Cr(VI) exposure monitoring. Our findings highlight distinct molecular trajectories in response to Cr(VI) over time, providing valuable insights into the progression from early toxic stress to chronic carcinogenic transformation. These results advance our understanding of Cr(VI)-induced carcinogenesis and suggest these potential targets for preventive and therapeutic interventions in exposed populations.

 $Keywords: hexavalent \ chromium; \ Cr(VI); \ transcriptomics; \ oxidative \ stress; \ carcinogenesis; \ biomarkers$

1. Introduction

Chromium (Cr) is an essential trace element in its trivalent form, Cr(III) but is highly toxic and carcinogenic in its hexavalent form Cr(VI) [1, 2].

Exposure to Cr(VI) is a well-documented occupational and environmental hazard, leading to severe health risks such as oxidative stress, DNA

tumorigenesis, and immune system dysfunction [3, 4]. Epidemiological studies have linked chronic Cr(VI) exposure to multiple adverse health outcomes. For instance, dermal exposure to Cr(VI) can cause skin irritation and ulceration [5], while workers in Cr(VI)-related industries experience a significantly higher risk of lung cancer compared with the general population [6]. Moreover, ingestion of drinking water or food contaminated with hexavalent chromium poses additional health risks [7]. The effects of Cr(VI) on the human body vary widely depending on exposure concentration, duration, and biological context. However, the comprehensive molecular characteristics distinguishing acute from chronic Cr(VI) toxicity remain poorly understood.

High-throughput transcriptomic analyses have provided valuable insights into the molecular pathways underlying Cr(VI)-induced toxicity. For example, Cr(VI) exposure in human dermal fibroblasts has been shown to enrich pathways involved in apoptosis and oxidative stress [8], while exposure of human bronchial epithelial cells promotes carcinogenic transformation through activation of oncogenic pathways [9]. In addition, the effects of oral Cr(VI) exposure have been investigated in cell culture as well as in mouse and rat models, revealing systemic toxicity [10, 11]. Recent research has also emphasized the roles of histone modifications and non-coding RNAs in Cr(VI)-induced carcinogenesis, highlighting the long-term epigenetic consequences of exposure [12, 13].

Despite extensive research, direct transcriptomic comparison between short-term and long-term Cr(VI) exposure models has not yet been conducted. This study was designed to address this gap. Because occupational Cr(VI) exposure commonly leads to lung cancer as well as irritation of the skin and airways [14], the present work focused on transcriptomic datasets related to these tissues. Two datasets from the Gene Expression Omnibus (GEO) database were selected for comparative analysis: GSE16349, representing short-term Cr(VI) exposure (16 hours) and profiling gene expression changes in human dermal fibroblasts [8]; and GSE24025, representing long-term exposure (4 weeks) capturing transcriptomic alterations Cr(VI)-transformed cell colonies [9]. This comparative analysis provides new insights into the temporal molecular responses to Cr(VI) toxicity and identifies potential biomarkers for occupational health risk assessment and early detection of Cr(VI)-induced diseases.

2. Materials and Methods

2.1 Data Collection and Selection Criteria

Two publicly available Gene Expression Omnibus (GEO) datasets from NCBI were selected to investigate the molecular responses to Cr(VI) exposure across different time scales. These datasets were chosen for their well-characterized experimental designs, relevance to Cr(VI) toxicity, and availability of high-quality gene expression profiles. The short-term dataset. GSE16349, contains microarray-based expression profiles from primary human dermal fibroblasts exposed to 5 µM Cr(VI) for 16 hours, capturing acute molecular responses [8]. The long-term dataset, GSE24025, includes expression profiles from immortalized epithelial cell colonies (BEAS-2B) subjected to chronic Cr(VI) exposure (0.5 µM for 4 weeks), allowing analysis of transcriptional adaptations under prolonged exposure [9]. Raw data were obtained as CEL files (GSE24025) or text-based tables (GSE16349), normalized, and prepared for downstream bioinformatics analysis using methods described in our previous studies [15-17].

2.2 Data Preprocessing and Quality Control

For GSE24025, raw CEL files were processed using the Affymetrix package in R. Robust Multi-array Average (RMA) normalization was applied to correct background noise and normalize gene expression intensities. Gene annotation was performed using the hugene10stprobesetSYMBOL function, ensuring accurate probe-to-gene mapping. The GSE16349 dataset, available in a pre-normalized format, was converted into log2-transformed values to maintain consistency across datasets. To evaluate data distribution and detect potential batch effects, boxplots and violin plots were generated for both datasets. Principal Component Analysis (PCA) was performed to assess clustering patterns between control and Cr(VI)-exposed groups. Hierarchical clustering analysis was conducted using the d3heatmap package, allowing visualization of gene expression differences across conditions. The dataset preprocessing and normalization procedures were conducted in R Studio (version 1.2.1335) with R version 4.0.3 following standard bioinformatics workflows [18-20].

2.3 Differential Gene Expression Analysis and DEG Selection

Expression matrices from GSE16349 (5 μ M Cr(VI), 16 h) and GSE24025 (0.5 μ M Cr(VI), 4 weeks) were processed and analyzed independently using the limma package. Probe-level signals were mapped to gene symbols, averaged across probes

corresponding to the same gene, and transformed to the log₂ scale. For GSE16349, a two-group comparison was performed between Cr(VI)-exposed and control samples. For GSE24025, comparisons were made between Cr large versus control and Cr small versus control groups. Moderated statistics were computed using empirical Bayes shrinkage. Genes were considered differentially expressed if they showed a fold change ≥ 1.2 with a Benjamini-Hochbergadjusted p < 0.05. In the long-term model, a "shared" gene set was defined as the intersection of the Cr_large and Cr_small DEG lists showing consistent directions of change. "Common DEGs" across exposure durations were defined as genes meeting the differential expression criteria in both datasets with concordant regulation.

2.4 Hierarchical Clustering and Principal Component Analysis

To assess sample distribution and transcriptional heterogeneity, hierarchical clustering analysis and PCA were performed. Heatmaps were generated using pheatmap in R to visualize gene expression patterns across control and Cr(VI)-treated groups. PCA plots were constructed using pca3d, allowing for three-dimensional visualization of sample clustering. The separation of Cr(VI)-exposed and control groups provided additional validation of dataset consistency and differential expression trends. The common DEGs were clustered and functionally annotated using the ClusterGVis R package (version 4.5.0) [21]. Normalized gene expression values were Z-score transformed and subjected to hierarchical clustering with default parameters to identify groups of genes exhibiting similar expression patterns. The resulting clusters were visualized as a heatmap using the ClusterGVis, which simultaneously displays gene expression profiles, cluster sizes, and average expression trends. Functional enrichment of each cluster was assessed using Gene Ontology Biological Process (GO:BP) and Kyoto Encyclopedia of Genes and Genomes (KEGG) analyses implemented in the clusterProfiler package (version 4.16.0) [22]. Enriched terms with adjusted p < 0.05 were considered significant and summarized alongside each gene cluster in the heatmap [23-26].

2.5 Pathway Enrichment Analysis

Biological processes associated with gene expression profiles were identified using MetaCore (GeneGo, St. Joseph, MN, USA). Analyses were performed on two gene sets: (i) the DEGs shared by short-term (GSE16349) and long-term (GSE24025) exposure to identify core pathways, and (ii) the DEGs specific to each dataset to capture duration-dependent

biology [27-30]. Statistical significance was assessed with two-tailed Fisher's exact tests (p < 0.05).

2.6 Protein-Protein Interaction (PPI) Network Analysis

To further elucidate the Cr(VI)-responsive molecular interaction networks, Protein-Protein Interaction (PPI) analysis was conducted using the **STRING** database (version https://string-db.org) [31]. A total of 250 DEGs were uploaded to STRING, with Homo sapiens selected as the reference organism. The minimum required interaction score was set to 0.7 (high confidence), and active interaction sources included experimental evidence, co-expression, and curated database annotations. K-means clustering (k = 10) was applied to partition the network into functionally distinct modules. The resulting interaction network was visualized in Cytoscape software (version 3.10.4) [32]. Network topological analysis was subsequently conducted using the cytoHubba plugin to identify hub genes based on three centrality measures: (1) degree centrality, representing the number of direct interactions; (2) betweenness centrality, indicating a node's control over information flow within the network; and (3) closeness centrality. Genes ranking within the top 10 for each measure were defined as hub genes [33-36].

3. Results

3.1 Differential Gene Expression Analysis Reveals Common and Distinct Responses to Short-term and Long-term Cr(VI) Exposure

The analysis workflow of this study is illustrated in Figure 1. To systematically compare transcriptomic alterations induced by short-term (16 h) and long-term (4-week) Cr(VI) exposure, we identified differentially expressed genes (DEGs) defined by fold change (FC) ≥ 1.2 and Benjamini-Hochberg-adjusted p < 0.05. In the short-term dataset (GSE16349), 5741 DEGs were detected (2825 upregulated; 2916 downregulated). In the long-term model (GSE24025), comparisons of Cr_large and Cr_small colonies versus untreated controls yielded 1590 and 2359 DEGs, respectively (Figure 2). Principal component analysis (PCA) revealed that Cr_large and Cr_small colonies exhibited highly similar transcriptional profiles (Supplementary Figure S1), which was further supported by hierarchical clustering analysis showing consistent gene expression patterns between the two groups (Supplementary Figure S2). Intersection analysis identified 1134 shared DEGs upregulated; 679 downregulated) between Cr_large and Cr_small colonies, justifying their combination into a unified long-term Cr(VI)-treated group for downstream analyses. To identify genes commonly responsive to Cr(VI) exposure across different durations, the 1134 shared DEGs from the long-term model were compared with those from the short-term dataset. This comparison revealed 86 genes consistently upregulated and 164 consistently downregulated, defining a core Cr(VI)-responsive signature conserved between acute and chronic exposure conditions.

To further explore distinct molecular responses under acute and chronic Cr(VI) exposure, clusterGVis analysis was performed to visualize gene clusters derived from GO and KEGG enrichment results. Specifically, 5741 DEGs from the short-term exposure

dataset (GSE16349) and 1134 DEGs from the long-term model (GSE24025) were subjected to functional enrichment and cluster visualization analyses, respectively, using the ClusterGVis package. In the short-term model (Figure 3), ClusterGVis identified four major gene clusters with distinct transcriptional patterns. Genes in clusters 1 and 2 were upregulated and enriched in biological processes related to DNA damage response, p53 signaling, and protein processing in the endoplasmic reticulum, reflecting acute stress responses. In contrast, clusters 3 and 4 were downregulated and primarily associated with cholesterol biosynthesis, metabolic regulation, and cell cycle progression.

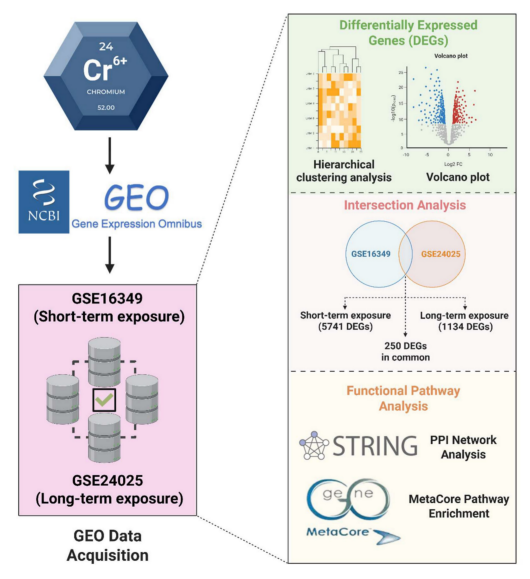


Figure 1. Overview of the study workflow. Short-term (GSE16349, 16 h) and long-term (GSE24025, 4 weeks) Cr(VI) exposure datasets were independently analyzed to identify DEGs. The intersection analysis defined shared DEGs and those unique to each exposure duration. Functional and network analyses were conducted using STRING for protein-protein interactions and MetaCore for pathway enrichment.

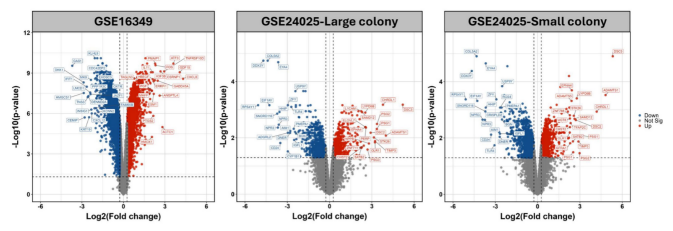


Figure 2. Volcano plots comparing chromium (VI)-exposed samples to matched controls. (A) GSE16349; (B) GSE24025-Large colony; (C) GSE24025-Small colony. Points are colored by significance: red, up-regulated; blue, down-regulated; grey, not significant. Vertical dashed lines mark |Fold change| = 1.2, and the horizontal dashed line marks p-value = 0.05). Top 20 DEGs are labeled.

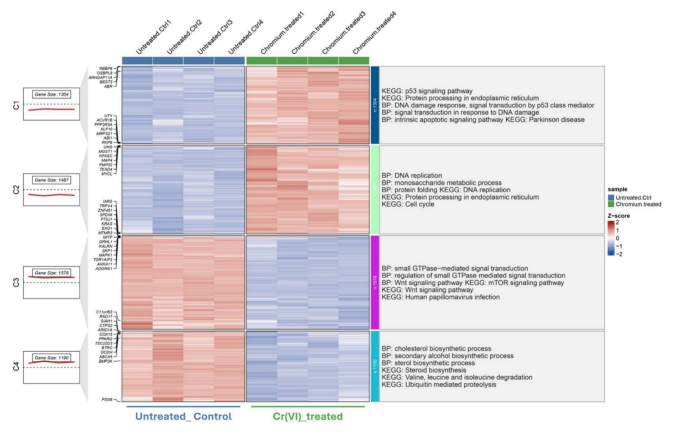


Figure 3. Transcriptomic clustering and functional enrichment of genes altered by Cr(VI) exposure (GSE16349). Hierarchical clustering of DEGs identified from GSE16349, comparing untreated control cells and Cr(VI)-treated cells. Each column represents an individual sample, and each row represents a gene. Expression values are shown as Z-scores, with red indicating upregulation and blue indicating downregulation relative to the mean expression. The left panels show the number of genes and average expression trend within each cluster. Functional enrichment analyses (Gene Ontology Biological Process and KEGG pathways) for each cluster are summarized on the right.

In the long-term model (Figure 4), five gene clusters were identified, highlighting persistent transcriptional reprogramming under prolonged Cr(VI) exposure. Upregulated clusters were significantly enriched in mitochondrial ATP synthesis, rRNA metabolic processes, and actin filament organization, whereas downregulated

clusters were associated with neuroactive ligand-receptor interaction and lipid metabolism. Collectively, these results indicate a shift from transient stress-related responses during short-term exposure to sustained metabolic and structural adaptation following chronic Cr(VI) treatment.

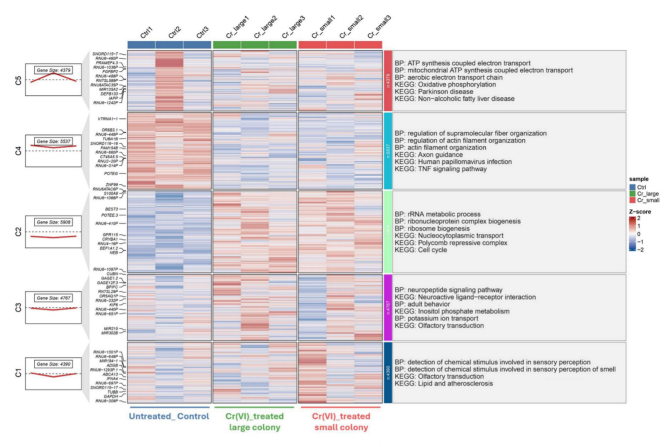


Figure 4. Transcriptomic clustering and functional enrichment of genes altered by Cr(VI) exposure (GSE24025). Hierarchical clustering of DEGs from dataset GSE24025, comparing untreated control (Ctrl), Cr(VI)-treated large colony, and Cr(VI)-treated small colony groups. Each column represents an individual sample, and each row represents a gene. Expression values are shown as Z-scores, with red indicating upregulation and blue indicating downregulation relative to the mean expression. The left panels show the number of genes and average expression trend within each cluster. Functional enrichment analyses (Gene Ontology Biological Process and KEGG pathways) for each cluster are summarized on the right.

3.2 Functional Pathway Enrichment Highlights Distinct Short-term and Long-term Cr(VI) Toxicity Mechanisms

MetaCore pathway enrichment analysis was applied to the common DEGs to identify core pathways activated under both short- and long-term exposure. Among the upregulated genes, the top enriched pathways were largely associated with apoptosis and survival signaling (nuclear PI3K/NGF-TrkA signaling, p53 activation, and APC-regulated cell cycle control), DNA damage response and repair (DNA replication initiation, mismatch repair, and apoptosis regulation), and cell cycle progression (initiation of mitosis and metaphase checkpoint). Additional categories included G-protein-mediated signaling (H-RAS, K-RAS, N-RAS, and Ga subunits), protein folding and maturation (insulin processing, CFTR regulation), and immune-related signaling (Treg regulation in COPD, IL-6 signaling in adipocytes, and CD4+ T-cell memory generation). Collectively, these findings suggest that Cr(VI) activates transcriptional exposure programs

promoting cell survival, stress adaptation, metabolic regulation, and immune modulation, consistent with pathways that support immune suppression and carcinogenic transformation (Figure 5, Supplementary Table S1). In contrast, the downregulated genes were enriched in pathways tied to cell cycle regulation and DNA damage checkpoints, such as ATM/ATRprogression, mediated G2/M control, G1/S prometaphase chromosome condensation, homologous recombination repair. Suppression was also observed in WNT/ β -catenin and TGF- β signaling (linked to hepatocellular carcinoma, lung cancer, and pancreatic cancer), pointing to impaired regulation of tissue homeostasis and tumor-suppressive signaling. Additional enrichment involved immune and inflammatory processes (IL-8 signaling, IFN- α/β -MAPK signaling, M-CSF receptor signaling, and Th17 differentiation) as well as fibrosis/ECM remodeling (stellate cell activation, Angiotensin II-PI3K/ERK signaling, and EMT regulation). Together, these results indicate that Cr(VI) exposure is associated with downregulation of genomic maintenance, checkpoint control, and immune surveillance, thereby

fostering genomic instability, immune evasion, and a tumor-permissive microenvironment (Figure 6, Supplementary Table S2).

further distinguish exposure-specific transcriptional responses, pathway enrichment analyses were performed on distinct DEGs from the acute (GSE16349) and chronic (GSE24025) models. In the acute exposure dataset (GSE16349), enriched pathways highlighted rapid activation of oxidative stress and hypoxia responses (ROS signaling, HIF-1 transcriptional targets, and negative regulation of HIF1A function). Robust engagement of DNA damage response and repair was also evident (ATM/ATR activation, p53 signaling, and DNA replication initiation/elongation), together apoptosis and survival pathways (p53p73-dependent apoptosis, PDGF-PI3K/AKT-NFkB, and mTORC1 signaling). Additional enriched categories included cytoskeletal remodeling and motility (Rho GTPase-regulated actin organization, receptor signaling, PDGF-Rho GTPase signaling), cell cycle checkpoints (Cullin1/Rbx1 E3 ligase control, G1/S transition, mitotic control), and oncogenic/developmental signaling (WNT/ β -catenin, NOTCH, TGF- β , and EGFR).

Immune-related and pro-fibrotic signaling (IL-3, IL-8, BCR, Th2/TNF-α-induced fibrosis, stellate cell activation) were also enriched. These results suggest that short-term Cr(VI) exposure triggers an immediate stress-response program characterized by ROS production, DNA damage repair, and apoptosis regulation, while simultaneously perturbing immune, developmental pathways. responses may serve as adaptive mechanisms to genotoxic stress but also create vulnerabilities that predispose carcinogenesis (Supplementary Figure S3, Supplementary Table S3). By contrast, the chronic exposure dataset (GSE24025) revealed sustained enrichment of fibrotic and pro-tumorigenic signaling pathways. These included stellate cell activation and liver fibrosis, multiple TGF-β-driven programs (stimulation in lung, breast, pancreatic, and colorectal cancers; SMAD signaling; Activin A signaling; and EMT induction), and EGFR- and WNT/β-cateninmediated transcriptional regulation. Chronic exposure also impacted cell cycle control, with enrichment of G1/S regulation, Cul1/Rbx1-mediated checkpoints, senescence, and metaphase transition. In addition, immune and inflammation-related pathways were prominent,

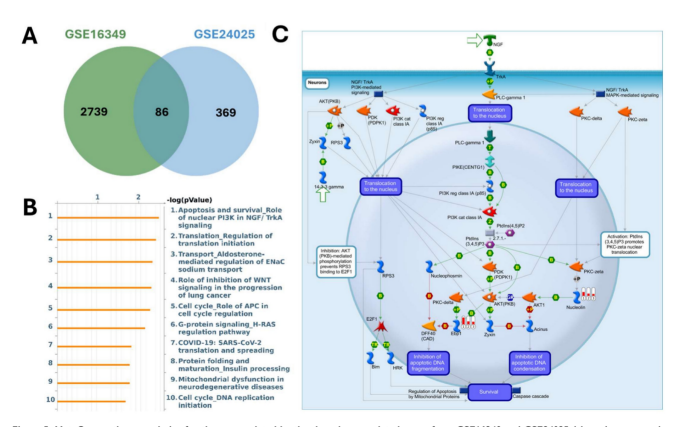


Figure 5. MetaCore pathway analysis of pathways regulated by the shared up-regulated genes from GSE16349 and GSE24025 (chromium-exposed vs control). (A) Venn diagram showing 2739 up-regulated genes unique to GSE16349, 369 genes unique to GSE24025, and 86 up-regulated genes in both datasets. (B) Top 10 enriched MetaCore pathways for the shared up-regulated set ranked by −log10(p-value). (C) MetaCore process map for the top pathway, "Apoptosis and survival-Role of nuclear PI3K in NGF/TrkA signaling," with the shared up-regulated genes overlaid, highlighting the PI3K/AKT-mediated survival module. Up-regulation was defined in each dataset as log2(FC) ≥ 1.2 with p-value < 0.05.

including IL-6 signaling (in breast and prostate cancer) and Hedgehog/IGF/HGF cooperation in stem cell regulation, suggesting acquisition of cancer stemness traits. ECM remodeling and fibrosis-related signatures (fibroblast/myofibroblast activation, systemic sclerosis, and metalloprotease signaling) further underscore the emergence of a fibrotic, tumor-permissive microenvironment (Supplementary Figure S4, Supplementary Table S4).

In summary, short-term exposure is dominated by acute stress responses (oxidative stress, DNA damage repair, and apoptosis), whereas long-term exposure reprograms transcription toward fibrosis, EMT, and oncogenic signaling. Together, these findings capture a continuum from immediate adaptation to chronic carcinogenic transformation.

3.3 Protein-Protein Interaction Network Identifies Key Hub Genes in Cr(VI) Exposure

Protein–protein interaction (PPI) analysis using the STRING database revealed a network comprising 250 nodes and 84 edges, indicating extensive interconnectivity among the DEGs (Figure 7). K-means clustering (k = 18) further partitioned the network into distinct submodules, many of which were enriched in ribosome biogenesis and protein folding, cell-cycle regulation, sterol biosynthesis, phosphatidylinositol phosphate metabolism, TGF- β activation, ubiquinol biosynthesis, and apoptosis-related pathways. The largest cluster contained AATF, CCT4, CCT5, CCT7, EIF3D, and

HMGN4, representing a tightly connected chaperonin-ribosomal module central to proteostatic control, while smaller clusters captured specialized metabolic and signaling processes such as LGALS3-mediated apoptosis and ferritin complex formation, underscoring the multifaceted cellular response to Cr(VI).

To identify key molecular mediators within this network, topological analysis was performed using the cytoHubba plugin in Cytoscape. Three centrality including degree, betweenness, closeness, were applied to evaluate node importance from complementary perspectives (Figure 8A-C). Across all three analyses, RPL27A, PA2G4, and PES1 consistently emerged as core hubs, underscoring their pivotal roles in maintaining network connectivity and mediating adaptation to chromium exposure. In contrast, CCT5, CCT7, and EIF3D ranked highly in two of the three analyses, reflecting strong but context-dependent participation in proteostasis and chaperonin-assisted folding. The degree and closeness networks were largely concordant, highlighting a compact nucleolar- translational core composed of RPL27A, PA2G4, PES1, and EIF3D, together with a CCT chaperonin mini-cluster (CCT5, CCT7). The betweenness map preserved this structure but additionally elevated AATF, SDAD1, and WDR46 as bridging nodes linking RNA-processing chaperone/cell-cycle modules.

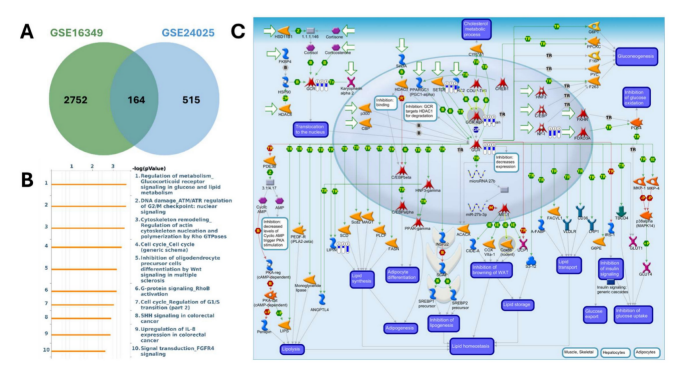


Figure 6. MetaCore pathway analysis of pathways regulated by the shared down-regulated genes from GSE16349 and GSE24025 (chromium-exposed vs control). (A) Venn diagram showing 2752 down-regulated genes unique to GSE16349, 515 genes unique to GSE24025, and 164 down-regulated genes in both datasets. (B) Top

10 enriched MetaCore pathways for the shared down-regulated set ranked by $-\log 10(p\text{-value})$; (C) MetaCore process map for the top pathway, "Regulation of metabolism-Glucocorticoid receptor signaling in glucose and lipid metabolism," with the shared down-regulated genes mapped, highlighting modules governing lipolysis, lipogenesis, adipogenesis, insulin signaling, and lipid homeostasis. Down-regulation was defined as $\log 2(FC) \le -1.2$ with p-value < 0.05.

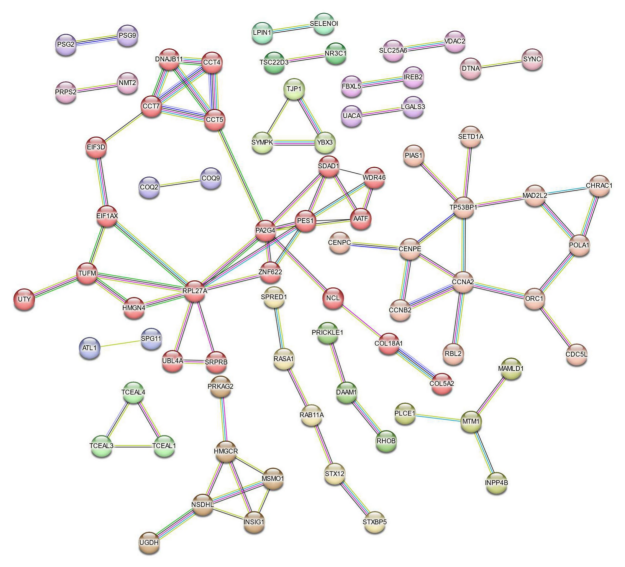


Figure 7. STRING-based Protein-Protein Interaction (PPI) network of Cr(VI)-responsive genes. The network consists of 84 edges (interactions) among 250 nodes (proteins) derived from Cr(VI)-responsive DEGs. A high-confidence interaction score (≥ 0.7) was applied, and nodes are color-coded by k-means clustering (k = 18). Only connected nodes are displayed.

When the top-ranked nodes from all three centrality metrics were integrated, six consensus hub genes, including RPL27A, PA2G4, PES1, CCT5, CCT7, and EIF3D, were identified, representing genes that combine extensive interactions (high degree and closeness) with critical information flow across functional (high submodules betweenness). Collectively, these findings define a tightly integrated PPI network organized around two principal functional axes: ribosome biogenesis and translation initiation (RPL27A, PA2G4, PES1, EIF3D) and proteostasis via the CCT foldase complex (CCT5, underscoring the nucleolar-ribosomal machinery as a central element of the cellular response to Cr(VI)-induced stress.

4. Discussion

study provides a comprehensive comparative transcriptomic analysis of short-term and long-term Cr(VI) exposure, revealing both shared and time-dependent molecular alterations. integrating two independent **GEO** representing acute (GSE16349, 16 hours) and chronic (GSE24025, 4 weeks) Cr(VI) exposure, we identified common DEGs and uncovered transcriptional programs underlying the transition from immediate stress responses to chronic carcinogenic transformation. A schematic summary of the findings is illustrated in Figure 9.

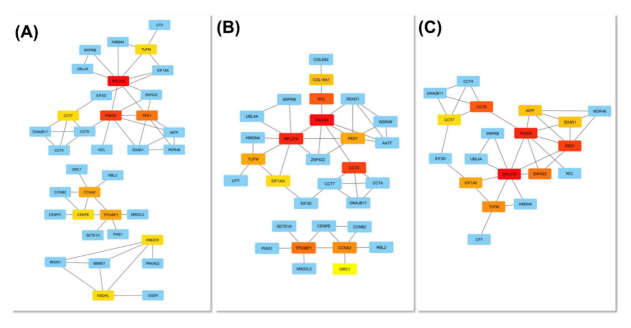


Figure 8. Identification of hub genes in the Cr(VI)-responsive PPI network. Hub genes were determined using three cytoHubba centrality metrics: (A) Degree, (B) Betweenness, and (C) Closeness. The top 10 ranked nodes for each metric are visualized in the corresponding subnetworks. Node color intensity represents hub ranking (red = higher centrality). Highly ranked nodes, particularly PL27A, PA2G4, and PES, formed a densely interconnected module enriched in ribosomal and nucleolar proteins, highlighting a nucleolar-ribosome-biogenesis axis as the topological core of the network.

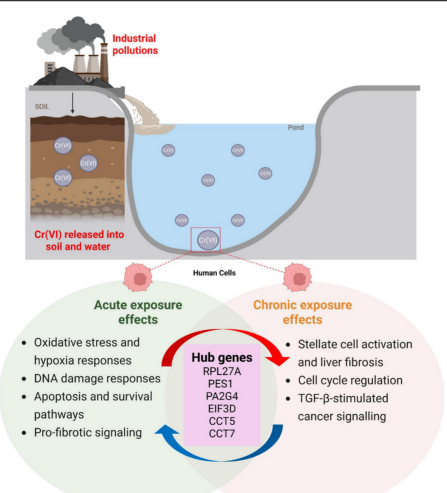


Figure 9. Proposed model summarizing the molecular effects of acute and chronic hexavalent chromium (Cr(VI)) exposure.

Our findings demonstrate that short-term Cr(VI) exposure primarily triggers acute stress responses characterized by activation of oxidative stress, DNA damage repair, and apoptotic pathways. This observation aligns with prior studies showing that Cr(VI) rapidly generates reactive oxygen species (ROS) and induces genotoxic stress, activating p53-dependent and p73-mediated apoptosis signaling [37-41]. The early upregulation of DNA repair and checkpoint genes observed here likely reflects a protective response aimed at maintaining genomic stability under transient oxidative insult. In contrast, prolonged exposure to Cr(VI) led to extensive transcriptional reprogramming associated fibrosis, epithelial-mesenchymal transition (EMT), and oncogenic signaling. Enrichment of TGF-β, WNT/β-catenin, and EGFR pathways in the chronic model suggests a shift toward pro-tumorigenic and pro-fibrotic signaling, consistent with the progression from adaptive to maladaptive responses. These results are consistent with previous findings that long-term Cr(VI) exposure promotes EMT, epigenetic alterations, and anchorage-independent growth in human bronchial epithelial cells [42-44]. Together, these observations suggest that sustained Cr(VI) exposure drives a temporal continuum of molecular events, from acute genotoxic stress to chronic transcriptional remodeling, culminating carcinogenic transformation.

PPI analysis highlighted network ribosomal-centered network dominated by RPL family members (RPL27A), along with nucleolar proteins such as PES1 and PA2G4. These hub genes are well-known regulators of ribosome biogenesis, translational control, and cell cycle progression. Notably, RPL5 and RPL11 form part of the 5S ribonucleoprotein complex that modulates p53 activity by sequestering MDM2, thereby linking ribosome biogenesis stress to genome surveillance [45, 46]. Their upregulation in both acute and chronic exposure models suggests persistent activation of nucleolar stress signaling. The concurrent enrichment of rRNA processing and mitochondrial translation pathways further supports the notion that Cr(VI) toxicity disrupts proteostasis and energy metabolism. Given that dysregulated ribosome biogenesis is increasingly recognized as a hallmark of cancer, the dominance of ribosomal proteins within the Cr(VI) PPI network underscores their potential role in driving chronic toxicity and malignant transformation [46-48]. Integrating functional enrichment results across both exposure models suggests a temporal shift in cellular priorities. During acute exposure, cells mount a coordinated antioxidant, DNA repair, and apoptotic response aimed at mitigating damage.

However, under sustained exposure, these defensive mechanisms give way to pathways promoting cell proliferation, ECM remodeling, and immune modulation. Suppression of tumor-suppressive signaling (WNT/ β -catenin and TGF- β regulation) and dysregulation of immune pathways (IL-6, IFN- α/β , and Th17 signaling) were observed, collectively favoring a microenvironment conducive to malignant transformation. These transcriptional transitions reflect the biological trajectory from oxidative stress-driven cytotoxicity to fibrotic remodeling and eventual neoplastic progression, consistent with epidemiological evidence linking chronic Cr(VI) exposure to lung carcinogenesis [49, 50].

The identification of common DEGs and hub genes across both exposure durations provides a foundation for developing biomarkers to monitor Cr(VI) exposure and early cellular transformation. In particular, RPL27A and PES1 may serve as sentinel indicators of Cr(VI)-induced nucleolar stress, while fibrosis-related genes involved in ECM remodeling and $TGF-\beta$ signaling could mark chronic adaptation and disease progression. Validation of these genes in exposed human cohorts or experimental animal models would strengthen their translational potential for occupational health surveillance.

Despite its integrative scope, this study has several limitations. First, the analysis relied on pre-existing datasets derived from distinct cell types and experimental conditions, which may introduce confounding biological variability. transcriptomic profiling captures only mRNA-level changes, while Cr(VI) toxicity also involves post-translational and epigenetic mechanisms not addressed here. **Future** studies combining transcriptomics with proteomic and epigenomic analyses will be essential to fully delineate the multilayered response to Cr(VI). Experimental validation of candidate biomarkers and hub genes in human tissues and in vivo models will further clarify their functional relevance and diagnostic utility.

5. Conclusion

conclusion, this study provides transcriptomic comprehensive comparison of responses to short-term and long-term hexavalent chromium exposure. Acute exposure primarily activated oxidative stress, DNA repair, and apoptotic pathways, whereas prolonged exposure promoted transcriptional reprogramming associated fibrosis, epithelial-mesenchymal transition, and oncogenic signaling. The consistent involvement of nucleolar and ribosomal proteins highlights their central role in stress adaptation and cellular transformation. Together, these findings offer new insights into the molecular progression from early toxic injury to chronic carcinogenesis and identify potential biomarkers for monitoring and mitigating Cr(VI)-induced health risks.

Supplementary Material

Supplementary figures and table. https://www.medsci.org/v23p0113s1.pdf

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Competing Interests

The authors have declared that no competing interest exists.

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