Knowledge Building Insights on Biomarkers of Arsenic Toxicity to Keratinocytes and Melanocytes

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Abstract: Exposure to inorganic arsenic induces skin cancer and abnormal pigmentation in susceptible humans. High-throughput gene transcription assays such as DNA microarrays allow for the identification of biological pathways affected by arsenic that lead to initiation and progression of skin cancer and abnormal pigmentation. The overall purpose of the reported research was to determine knowledge building insights on biomarker genes for arsenic toxicity to human epidermal cells by integrating a collection of gene lists annotated with biological information. The information sets included toxicogenomics gene-chemical interaction; enzymes encoded in the human genome; enriched biological information associated with genes; environmentally relevant gene sequence variation; and effects of non-synonymous single nucleotide polymorphisms (SNPs) on protein function. Molecular network construction for arsenic upregulated genes \textit{TNFSF18} (tumor necrosis factor [ligand] superfamily member 18) and \textit{IL1R2} (interleukin 1 Receptor, type 2) revealed subnetwork interconnections to \textit{E2F4}, an oncogenic transcription factor, predominantly expressed at the onset of keratinocyte differentiation. Visual analytics integration of gene information sources helped identify \textit{RAC1}, a GTP binding protein, and \textit{TFRC}, an iron uptake protein as prioritized arsenic-perturbed protein targets for biological processes leading to skin hyperpigmentation. \textit{RAC1} regulates the formation of dendrites that transfer melanin from melanocytes to neighboring keratinocytes. Increased melanocyte dendricity is correlated with hyperpigmentation. \textit{TFRC} is a key determinant of the amount and location of iron in the epidermis. Aberrant \textit{TFRC} expression could impair cutaneous iron metabolism leading to abnormal pigmentation seen in some humans exposed to arsenicals. The reported findings contribute to insights on how arsenic could impair the function of genes and biological pathways in epidermal cells. Finally, we developed visual analytics resources to facilitate further exploration of the information and knowledge building insights on arsenic toxicity to human epidermal keratinocytes and melanocytes.

Keywords: arsenic, disulfide bond, environmental response genes, functional annotation, hyperpigmentation, iron uptake, keratinocyte, melanocyte, RAC1, SNPs, skin cancer, TFRC, toxicogenomics, vicinal cysteines

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Introduction
Arsenic is a naturally occurring environmental toxicant that is widely distributed in the earth’s crust.\(^1,2\) The air, water, and soil environment can be contaminated with arsenic from industrial smelting of metals, power generation with coal, and applications of pesticides and herbicides.\(^3\) Human exposure occurs mainly through the accumulation of arsenic from soil into food sources including grains, vegetables, fish, and meats.\(^4\,6\) Arsenic enters the human body by dermal contact, inhalation, or food consumption, but ingestion of contaminated drinking water is the most common route. The tissues most often affected by arsenic exposure are the skin, nasal passages, lungs, gastrointestinal tract, and liver.\(^7\)

Human exposure to chronic low-dose arsenic is associated with an increased risk of epithelial cancers of the skin such as intraepidermal carcinomas (Bowen disease), squamous cell carcinomas (SCC), basal cell carcinomas (BCC),\(^8\) and Merkel cell carcinoma (MCC).\(^9\) In most cases where internal cancers are attributed to arsenic exposure, there are other evidences of cutaneous cancers due to the adverse effects of arsenic expressed in the form of arsenical keratosis, hyperpigmentation, and multiple cutaneous malignancies.\(^10\) Clinical manifestations of arsenic exposure include epithelial cancers of the bladder, liver, kidney, prostate, and lung; cardiovascular and skin diseases; as well as neuropathies of the central nervous system.\(^7\)

The mechanisms of toxicity and carcinogenicity of arsenicals on various cell types have been proposed.\(^11\,15\) These mechanisms include perturbation of signaling cascades, oxidative stress, and chromosomal aberrations.\(^16\) There could also be perturbation of the transcriptional activity and changes in the global gene expression.\(^15\,17\,18\) Exposure to arsenic trioxide leads to the alteration in mitochondrial integrity and the generation of reactive oxygen species (ROS),\(^19\) enhanced cell proliferation,\(^20\) and alteration of DNA methylation.\(^21\)

Advances in genomics and other “-omics” technologies are providing massive numbers of datasets and the accompanying scientific publications that describe potential gene, protein, and biological processes as potential biomarkers of adverse health effects of environmental chemicals. In particular toxicogenomics, which is a scientific field that investigates how the entire genome response to environmental toxicant effects, is a recognized approach to discover potential biomarkers of adverse toxicity and exposure as well as to validate/quantify biomarker signatures.\(^22\) The growth in data from toxicogenomics research has led to the development of bioinformatics databases such as the Comparative Toxicogenomics Database (CTD) for curating toxicogenomics relationships (chemical-gene, chemical-disease, and gene-disease) found in scientific publications.\(^23\) As of the December 7, 2011, the CTD contained 28,413 PubMed references, 352,925 chemical-gene relationships, 6,605 unique chemicals, 20,710 unique genes, and 334 unique organisms. These toxicogenomics relationships and data when combined with biological information on human genes from other bioinformatics databases can lead to knowledge building (discovering previously unknown relationships from datasets) on potential biomarkers. Significant over-representation (enrichment) of certain biological topics for a particular gene is an example of the biological information that can be obtained from a bioinformatics database such as ConceptGen.\(^24\) ConceptGen database currently consists of approximately 18,000 concepts, each with 5 or more assigned genes. Further, diverse bioinformatics tools are now available to reconstruct molecular interactions from predicted and experimentally validated datasets.\(^25\,29\) The reconstruction of molecular pathways involving potential biomarker genes and proteins can also yield insights on how the normal cellular activities are altered in different chemical exposures or disease conditions. These diverse datasets from bioinformatics databases, therefore, present opportunities for discovery and inferences on the biological processes affected by arsenicals. In particular, we are interested in identifying genes for further research on the mechanisms of toxic action and cancer initiation of arsenic on skin keratinocytes.

We have used a visual analytics approach to integrate and make sense of data on arsenic-associated genes stored in two bioinformatics resources: Comparative Toxicogenomics Database and ConceptGen. According to Boulos et al,\(^30\) the goal of visual analytics is to facilitate the discourse between the user and the data by providing dynamic displays and versatile visual interaction opportunities with the data that can support analytical reasoning and the
exploration of data from multiple user-customizable aspects. Interactive visual analytics approaches are increasingly required to assist human cognition and analytical reasoning to identify patterns in datasets of diverse sizes. The primary objectives of the reported research were to (1) compare the toxicogenomics relationships for genes in a molecular network involving two genes (IL1R2 and TNFSF18) that were upregulated in HaCaT keratinocytes exposed to chronic, low dose arsenic trioxide; and (2) identify arsenic-interacting genes enriched for the melanosome (melanin-pigment bearing organelles transferred from melanocytes to keratinocytes) biological information. These objectives are to help generate hypotheses for further research on the mechanisms of arsenic toxicity on skin keratinocytes, in particular, aberrant skin pigmentation and cancer initiation.

Our visual analytics protocols identified (1) five genes in the IL1R2 and TNFSF18 molecular network that were perturbed by an arsenical in at least one of the three keratinocyte cell lines (2) 106 genes were enriched for the skin pigmentation topics: melanocyte, melanogenesis, melanosome(s), melanocyte differentiation, melanosome membrane, and skin pigmentation and (3) nine genes were localized in the melanosome but also known to be perturbed by arsenic trioxide in other cell types. Molecular network construction on selected upregulated genes encoding membrane proteins revealed interconnections to E2F4, an oncogenic transcription factor, predominantly expressed at the onset of keratinocyte differentiation. Two melanosome localized proteins TFRC and RAC1 were also prioritized from the visual analytics process. The reported findings provide knowledge building insights on how arsenic could impair the function of genes and the biological pathways related to arsenic toxicity in keratinocytes and melanocytes.

Methods
Data sources for knowledge building insights on arsenic toxicity to keratinocytes and melanocytes
The data sources for gene lists used in the research were (1) molecular interaction of arsenic-regulated genes IL1R2 and TNFSF18, (2) Comparative Toxicogenomics Database, (3) ConceptGen database, (4) Human Genome Organization Nomenclature Committee Gene Names, (5) SNPs3D (molecular functional effects of non-synonymous SNPs based on structure and sequence analysis), and (6) Environmental Genome Sequencing Project. The retrieval and processing of the gene lists are described in the following sections.

Prediction of molecular network of genes for arsenic-regulated genes IL1R2 and TNFSF18
Genes for interleukin 1 receptor, type II (IL1R2) and tumor necrosis factor (ligand) superfamily, member 18 (TNFSF18) encode membrane proteins that form ligand-receptor systems. We therefore assumed that the predicted molecular interactions for IL1R2 will have connections to ligands IL1B and IL1A, while TNFSF18 will have connection to its receptor TNFRSF18. We used the Michigan Molecular Interactions (MiMI) plugin for Cytoscape to retrieve molecular interactions as well as interaction attributes in the Michigan Molecular Interactions resource for query genes IL1B, IL1R2, IL1A, TNFSF18, and their nearest neighbors. The Cytoscape platform was used to display the biological networks with the biological entities (genes/proteins) represented as nodes and the biological interactions represented as edges between nodes.

Construction of arsenical-gene interactions datasets
Arsenicals are defined in the Medical Subject Headings database (http://www.ncbi.nlm.nih.gov/mesh) as inorganic or organic compounds that contain arsenic, and the term is assigned an accession number of D001152. The Batch Query tool (http://ctdbase.org/tools/batchQuery.go) of the Comparative Toxicogenomics Database was used to download the arsenical-gene interaction datasets for (1) all the genes curated in the Comparative Toxicogenomics Database having at least an interaction with arsenicals and (2) all the genes in the predicted molecular network involving IL1R2 and TNFSF18. In both cases, the fields that constitute each record were Input (Gene Symbol or NCBI Entrez Gene ID), GeneSymbol (Approved Gene Symbol), GeneName (Approved Gene Name), GeneID (Entrez Gene ID), ChemicalName (NCBI MeSH Chemical Name), ChemicalID (NCBI MeSH Chemical ID), CasRN (Chemical Abstracts Service Number), and ChemicalFormula.
Registry Number), Organism (Scientific Name of Organism), OrganismID (NCBI Taxonomy ID), Interaction (Curated Interaction), InteractionActions (Effect of chemical on gene), and PubMedIDs (PubMed Identifier). The CTD chemical-gene interaction data were determined from peer-reviewed articles published and indexed in PubMed (www.pubmed.gov).

For the molecular interaction network, each record of the dataset was further annotated with the cell or organ type, sequence type (mRNA or protein), chemical classification (arsenical or non-arsenical) and the in vivo or in vitro experimental systems associated with the entry. The additional annotations (data fields) were determined after reading the abstract and full text of the PubMed indexed article. The sequence type category was included to identify proteins that may bind to arsenic via vicinal cysteines.

Construction of dataset on enriched biological information for arsenic-interacting genes
Enriched biological topics for genes curated to interact with arsenicals were programatically retrieved from the ConceptGen data repository. Each gene can be enriched in 14 concept types: Gene Ontology (GO) biological process; GO molecular function; GO cellular component; Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway; Biocarta pathway; Protein ANalysis THrough Evolutionary Relationships (Panther) Pathways; Protein Domains (Pfam); Medical Subject Headings (MeSH); Online Mendelian Inheritance in Man (OMIM); Drug Bank Target Sets; microRNA Predicted Tagets; Gene Expression Omnibus (GEO) datasets, Michigan Molecular Interactions (MiMI) and Protein Interactions; Metabolites; and Cytoband (chromosomal locations). The dataset constructed consists of Gene Symbol, ConceptID, Concept Name and Concept Type.

Dataset of human genes encoding enzymes
Since enzymes are components of biological pathways, we were interested in determining the subset of CTD arsenic-interacting genes that are enzymes. The complete Human Genome Nomenclature Committee (HGNC) gene names dataset consisting of approved symbols of over 33,000 loci were downloaded from www.genenames.org.

Dataset on prioritized environmental response human genes
The gene symbols for 647 prioritized environmental response human genes were obtained from the National Institute of Environmental Health Sciences’ Environmental Genome Project (NIEHS EGP) at http://egp.gs.washington.edu/. These genes are involved in DNA repair, cell cycle regulation, apoptosis, and metabolism and are suspected to have a function in the susceptibility to environmental exposures in a panel of 95 individuals representing the ethnic diversity found in the United States. The putative functional effects of non-synonymous coding single nucleotide polymorphisms (ncSNPs) on the prioritized genes were extracted from NIEHS EGP website using a set of computer scripts. Two programs SIFT (Sorting Intolerant from Tolerant) and Polyphen (polymorphism phenotyping) were used to classify the ncSNPs. The SIFT classifies ncSNPs into tolerant or intolerant, while Polyphen classifies ncSNPs to benign, possibly damaging, and probably damaging. We were interested in genes with ncSNPs that change cysteine residues in the protein sequence. These ncSNPs may help to identify proteins that have vicinal cysteines, which are possible targets for arsenic binding.

Dataset of arsenic-interacting genes with candidate SNPs affecting disulfide bonds
The supplementary table from our previous publication on candidate single nucleotide polymorphisms for arsenic responsiveness in proteins was retrieved from the PubMed Central website for the article (http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2964045/bin/BBI-4-supplementary.xls). The set of genes that have breakage of disulfide bond as annotation of the impact of SNP on protein stability was selected for further analysis.

Visual analytics of datasets (gene lists)
The GeneSymbol (Gene Symbol) field was present in all the datasets analyzed and, therefore, was used to link the datasets. A visual analytics software, Tableau Professional (http://tableausoftware.com/), was used to address the primary research objectives: (1) compare the toxicogenomics relationships for genes in a molecular network involving the two genes (IL1R2 and TNFSF18) that were upregulated in HaCaT keratinocytes exposed to chronic, low dose arsenic.
Knowledge building insights on arsenic toxicity in skin cells

Background

Trioxide and (2) identify arsenic-interacting genes enriched for the melanosome topics (melanin-pigment bearing organelles transferred from melanocytes to keratinocytes). Additional objectives conducted to support the two primary objectives were to (1) identify the arsenic-interacting enzymes with melanosome localization that are perturbed by arsenic; (2) identify the environmental response human genes with potential vicinal cysteines for arsenic binding; and (3) design a visual analytic dashboard to enable integration and interactive exploration of data on effects of arsenic on genes with candidate single nucleotide polymorphisms affecting disulfide bonds. In each case, the data fields in the datasets were arranged on the visual analysis interface to enable interaction, integration, and presentation of the contents of the data fields. For example, in the integration of the CTD and ConceptGen datasets (gene lists), the following data fields were designed to be listed in the following order: GeneSymbol (CTD), Concept Name (ConceptGen), Interaction (CTD), Concept Type (ConceptGen), ChemicalName (CTD), and PubMedIDs (CTD) (Fig. 1).

Results

Data sources for knowledge building insights

The data sources for developing knowledge building insights using a visual analytics approach are presented in Table 1. The starting data source was a list of 3,826 genes curated to interact with arsenicals in the Comparative Toxicogenomics Database. The ConceptGen data was available for 3,419 arsenic-interacting genes. Additionally, 647 environmental response genes from the Environmental Genome Project were retrieved as well as 79 genes with associated SNPs that target cysteine residues. Finally, 36,805 human gene names were obtained from the Human Genome Organization Nomenclature Committee (HGNC) website (www.genenames.org). Table 2 is a list of 14 genes that were found in all the data sources. We integrated the gene list with Gene Ontology (GO) Cellular Component annotation stored in ConceptGen to determine those found in skin cell types. The RAC1, Ras-related C3 botulinum toxin substrate 1, met our filtering criteria as encoding a protein localized to the melanosome. The RAC1-arsenic interactions and associated PubMed Identifiers of the source articles are presented in Table 3. Visual analytics workbook is available as a Supplemental File and can be viewed with the free Tableau Reader (http://www.tableausoftware.com/products/reader).

Genes in molecular network perturbed by arsenicals in keratinocyte cell lines

A 24-node molecular interaction network containing IL1R2 and TNFSP18 was predicted using the MiMI
Table 1. Data sources and data types for knowledge building on arsenic toxicity.

| Data sources and data types                                                                 | Count  |
|---------------------------------------------------------------------------------------------|--------|
| Comparative toxicogenomics database (http://ctdbase.org/)                                  |        |
| Arsenic interacting genes                                                                  | 3,286  |
| Arsenicals                                                                                 | 29     |
| Arsenical-gene interactions                                                                 | 7,724  |
| Arsenicals interaction actions                                                              | 207    |
| Organisms investigated for arsenical action                                                 | 20     |
| PubMed identifiers for articles on arsenic interacting genes                                | 591    |
| ConceptGen (http://conceptgen.ncibi.org/)                                                  |        |
| Arsenic interacting genes in conceptGen                                                    | 3,419  |
| Concept names for arsenic interacting genes                                                | 13,587 |
| Concept types for arsenic interacting genes:                                               |        |
| Biocarta pathway                                                                            | 3,486  |
| Cytoband                                                                                   | 5,192  |
| Drug bank                                                                                  | 1,387  |
| Gene expression                                                                             | 19,2432|
| GO biological process                                                                       | 70,403 |
| GO cellular component                                                                      | 14,286 |
| GO molecular function                                                                       | 25,266 |
| KEGG pathway                                                                               | 6,198  |
| MeSH                                                                                       | 73,530 |
| Metabolite                                                                                 | 7,315  |
| miRBase                                                                                    | 12,8227|
| OMIM                                                                                       | 252    |
| Panther pathway                                                                             | 2,526  |
| pFAM                                                                                       | 6,083  |
| TransFac                                                                                   | 1,035  |
| Environmental genome project (http://egp.gs.washington.edu/)                               | 647    |
| Arsenical interacting genes with single nucleotide polymorphisms targeting cysteine residues | 79     |
| Human gene names (http://www.genenames.org)                                                | 36,805 |

Cytoscape Plugin (Fig. 2). As expected, TNFSF18 interacted directly with its receptor, TNFRSF18. Further, IL1R2 interacted directly with its ligands, IL1A and IL1B. An oncogenic transcription factor, E2F4, served as connecting node for the subnetwork containing the interleukin genes and the subnetwork containing TNFSF18 and TNFRSF18. The molecular interaction map also predicted Neddin (NDN) as a connection between E2F4 and IL1A. TNF receptor associated factor 2 (TRAF2) was predicted as a connection between E2F4 and TNFSF18. The activating transcription factor-2 (ATF2), a sequence-specific DNA-binding protein, was predicted to interact with both E2F4 and IL1B.

The clustering coefficient for the network of 24 genes was 0.128, which is a factor of the connectivity of a node and the connectivity of the neighborhood to which this node is connected. The average number of neighbors was 2.833. Another parameter examined was the connection degree, which measures the number of partners directly connected to a particular node. The connection degrees for IL1A, IL1B, IL1R2, TNFRSF18 and TNFSF18 were 9, 7, 4, 4, and 3 respectively.

The dataset in the CTD for the predicted molecular network consisted of 2,792 chemical-gene interactions (as of August 10, 2011) for 16 of the 24 genes. Figure 2 is a dashboard view consisting of 4 views generated from the dataset. These interactions, which consist of 676 chemicals investigated in 22 organisms, were divided into arsenical and non-arsenical groups. A set of 16 interactions were not annotated with organism information. All genes except S100 A13 (S100 calcium binding protein A13) had at least a curated interaction with an arsenical. The expression of five genes ATF2, CASP1, IL1A, IL1R2, and TNFSF18 were shown to be perturbed by arsenicals in the following keratinocyte cell lines: immortalize...
Table 2. Genes observed in all the five gene information sources for knowledge building insights on arsenic toxicity to keratinocytes and melanocytes.

| Gene symbol | Approved name                                                                 | CTD | ConceptGen | EGP | HGNC | SNPs3D |
|-------------|--------------------------------------------------------------------------------|-----|------------|-----|------|--------|
| ABCC1       | ATP-binding cassette, sub-family C (CFTR/MPR), member 1                       | 28  | 309        | 1   | 1    | 13     |
| ATM         | Ataxia telangiectasia mutated                                                 | 8   | 233        | 1   | 1    | 1      |
| BRCA1       | Breast cancer 1, early onset                                                  | 4   | 316        | 1   | 1    | 3      |
| BRCA2       | Breast cancer 2, early onset                                                  | 3   | 314        | 1   | 1    | 1      |
| CASP2       | Caspase 2, apoptosis-related cysteine peptidase                              | 4   | 72         | 1   | 1    | 1      |
| CDH1        | Cadherin 1, type 1, E-cadherin (epithelial)                                  | 7   | 1267       | 1   | 1    | 1      |
| EEF1A1      | Eukaryotic translation elongation factor 1 alpha 1                           | 3   | 140        | 1   | 1    | 1      |
| ERCC5       | Excision repair cross-complementing rodent repair deficiency, complementation group 5 | 3   | 119        | 1   | 1    | 1      |
| IGFBP7      | Insulin-like growth factor binding protein 7                                  | 2   | 106        | 1   | 1    | 1      |
| LTA         | Lymphotoxin alpha (TNF superfamily, member 1)                                | 4   | 471        | 1   | 1    | 1      |
| MAPK9       | Mitogen-activated protein kinase 9                                            | 26  | 160        | 1   | 1    | 2      |
| MSH5        | Muts homolog 5 (E. coli)                                                     | 1   | 109        | 1   | 1    | 1      |
| RAC1        | Ras-related C3 botulinum toxin substrate 1 (rho family, small GTP binding protein Rac1) | 11  | 339        | 1   | 1    | 3      |
| XPC         | Xeroderma pigmentosum, complementation group C                              | 4   | 120        | 1   | 1    | 1      |

Notes: CTD: Comparative Toxicogenomics Database (http://ctdbase.org/); ConceptGen: ConceptGen database at National Center for Integrative Biomedical Informatics (http://conceptgen.ncibi.org/); EGP: Environmental Genome Project Gene-Oriented Sequence Variation Discovery (http://egp.gs.washington.edu/); HGNC: Human Genome Organization Gene Nomenclature Committee (http://www.genenames.org/); SNPs3D: Molecular functional effects of non-synonymous SNPs based on structure and sequence analysis (http://www.snps3d.org/).

Table 3. Interactions of arsenicals with RAC1 protein and mRNA.

| Interaction                                                                 | PubMed IDs |
|-----------------------------------------------------------------------------|------------|
| [Ethanol co-treated with sodium arsenite] results in increased localization of RAC1 protein | 18182400   |
| [Tetradecanoylphorbol acetate co-treated with arsenic trioxide] results in increased expression of RAC1 mRNA | 18922899   |
| [Tetradecanoylphorbol acetate co-treated with arsenic trioxide] results in increased expression of RAC1 protein | 18922899   |
| NSC 23766 inhibits the reaction [sodium arsenite results in increased activity of RAC1 protein] | 19349368   |
| Pertussis Toxin inhibits the reaction [sodium arsenite results in increased activity of RAC1 protein] | 19349368   |
| RAC1 protein affects the reaction [sodium arsenite results in increased expression of PECAM1 protein] | 19033667   |
| S1PR1 protein affects the reaction [sodium arsenite results in increased activity of RAC1 protein] | 19349368   |
| Sodium arsenate results in increased expression of RAC1 mRNA                | 16882884   |
| Sodium arsenite results in increased activity of RAC1 protein               | 19349368   |
| Sodium arsenite results in increased localization of and results in increased activity of RAC1 protein | 18182400   |
| Sodium arsenite results in increased localization of RAC1 protein          | 18182400   |

Notes: *Abstract of article can be obtained from www.pubmed.gov. Additional information on arsenic-gene interactions can be obtained from http://ctdbase.org.
activity known to be perturbed by arsenic trioxide and other arsenicals in other cell types (Fig. 3). These gene-arsenic interactions were documented in 14 PubMed indexed articles (PubMed Identifiers shown in Fig. 3). We further prioritized these 22 genes based on the cell type used in the investigation with particular interest on skin cell types. The gene encoding an iron uptake protein, transferrin receptor (TFRC), was among the genes enriched for melanosome topic. The expression of TFRC was increased

Figure 2. Integrated view of chemical-gene interactions for molecular network with arsenic-upregulated genes TNFSF18 and IL1R2. Notes: Interaction Map for IL1B, IL1R2, IL1A, TNFRSF18 and TNFSF18 reveals E2F4 Transcription Factor as link between subnetworks. E2F4 controls cell cycle and acts on tumor suppressor proteins. The expression of five genes ATF2, CASP1, IL1A, IL1R2 and TNFSF18 were shown to be perturbed by arsenicals in keratinocyte cell lines: immortalize human keratinocytes (HaCaT), murine keratinocyte cell line (HEL30) and normal human epidermal keratinocyte (NHEK). Visual Analytics resource for exploring and downloading views is available at http://public.tableausoftware.com/views/tnfsf18_il1r2_ctd/molnet_ctd.

Figure 3. Selected arsenic-interacting genes enriched for melanosome topics. Notes: A total of 106 arsenic-interacting genes were enriched for the following skin pigmentation topics in ConceptGen. A subset of 22 genes is shown that were annotated to encode products that localize to the melanosome but their expression is known to be perturbed by arsenic trioxide and other arsenicals in other cell types. The changes in expression is limited to increase or decrease expression.
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by sodium arsenite in normal human epidermal keratinocytes (NHEK).13

**Arsenic-perturbed genes encoding enzymes localized in the melanosome**

Integration of ConceptGen and HGNC datasets identified 8 enzymes localized in the melanosome and were also observed to be perturbed by arsenic in other cell types (Fig. 4). The enzymes were carbonic anhydrase II (CA2), cathepsin B (CTSB), cathepsin D (CTSD), glucosidase, alpha; neutral AB (GANAB), matrix metallopeptidase 1 (interstitial collagenase) (MMP1) and protein disulfide isomerase family A, member 4 (PDIA4). The gene lists integration revealed in arsenic trioxide and sodium arsenite had a decreased expression of MMP1 mRNA in leukemia and keratinocytes cell lines respectively.13,41 Further, the PDIA4 mRNA had an increased expression from sodium arsenite in a leukemia cell line but a decreased expression by tetraarsenic tetrathiol in human vascular endothelial cells.42,43 Interaction of the two cathepsins (CTSB, CTSD) with arsenicals led to an increased protein or transcription levels.

**Environmental response genes with potential vicinal cysteines for arsenic binding**

The visual analytics pipeline retrieved 17 genes encoding protein isoforms that could have vicinal cysteines liable to arsenic binding. The gene for hemachromatosis (iron overload) was present in this list of genes. Molecular interaction network construction using MiMi cytoscope plug-in revealed the HFE interacts with Transferrin Receptor (TFRC), Transferrin Receptor 2 (TFR2) and Beta-2-Microglobulin (B2M) (Fig. 5).

**Arsenic-gene interaction for skin keratinocyte relevant genes with candidate SNPs targeting disulfide bonds**

A total of six genes (FST, HLA-C, ILA, KLK7, TFRC, and TLR4) were identified as having single nucleotide polymorphisms that cause breakage of disulfide bond. The visual analytics dashboard was designed to facilitate knowledge building insights on arsenic toxicity (Fig. 6). The dashboard integrates data streams from

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**Figure 4.** Enzymes localized in the melanosome and observed to be perturbed by arsenic in other cell types.

**Notes:** The enzymes lists were obtained by integrating datasets from ConceptGen and the Human Genome Organization (HUGO) Nomenclature Committee’s gene names website (http://www.genenames.org/). CA2: carbonic anhydrase II, CTSB: cathepsin B, CTSD: cathepsin D, GANAB: glucosidase, alpha; neutral AB, matrix; MMP1: metallopeptidase 1 (interstitial collagenase) and PDIA4: protein disulfide isomerase family A, member 4.
Figure 5. Integration of gene information sources for selected environmental response genes that are potential arsenic-binding proteins. Notes: The list of arsenic response genes was obtained from the Environmental Genome Project (http://egp.gs.washington.edu/) and the approved names from the Human Genome Organization Nomenclature Committee (HGNC) website (www.genenames.org). Arsenic-protein interaction relationships were obtained from Comparative Toxicogenomics (CTD http://ctdbase.org/). Molecular interaction of HFE gene was obtained from Michigan Molecular Interaction (MiMI). The color-coded boxes indicate the type of interaction with arsenic. The position of each box corresponds to the position of the word “protein” in the interaction text from CTD.

Figure 6. Visual analytics resource for exploring arsenic-interacting genes with candidate single nucleotide polymorphisms affecting disulfide bonds. Notes: The dashboard view enables user to search SNPs3D (www.snps3d.org) and PubMed (www.pubmed.gov) for genes with SNP associated breakage of disulfide bonds. Additional impacts on protein stability and genes available in as supplementary file of previous publication: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2964045/bin/BBI-4-supplementary.xls.
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Discussion

We report, for the first time, a unique visual analytics approach to integrate and interactively explore the gene information sources to determine knowledge building insights on arsenic-induced hyperpigmentation and skin cancer. The overall purpose of the reported research was to determine knowledge building insights on biomarker genes for arsenic toxicity to human epidermal cells by integrating a collection of gene lists annotated with biological information.

The information sets included toxicogenomics gene-chemical interaction, enzymes encoded in the human genome, enriched biological information associated with genes, environmentally relevant sequence variation, and effects of non-synonymous Single Nucleotide Polymorphisms on protein function.

Two datasets on arsenic-gene interactions form the basis of the reported research. First, we constructed a dataset of chemical-gene interactions for a network of 16 genes that included two genes upregulated in HaCaT keratinocytes exposed to chronic (22 passages), low-dose (0.5 mg/L) arsenic trioxide. Additionally, we enriched the dataset with data on cell/organ type and in vitro/in vivo experimental system for each chemical-gene interaction (Fig. 2). Second, the arsenic-gene interaction dataset of 3,826 genes curated for interaction with arsenic was integrated with other datasets for knowledge building insights on arsenic-gene interaction in the melanosome. The human epidermal melanosomes are pigment (melanin)-containing large elliptical or spheroidal organelles (approximately 500 nm in diameter) formed in the melanocytes, donated to neighboring keratinocytes by passing through dendrites of melanocytes. Taken together, the two starting datasets, the additional annotations, the data sources, and the visual analytics approach enabled a set of research objectives to be accomplished.

We have previously reported that the genes for cytokines TNFSF18 and ILIR2 are perturbed when human skin keratinocytes (HaCaT) was chronically exposed to low dose of arsenic trioxide. TNFSF18 is a ligand for receptor TNFRSF18 and it modulates T-lymphocyte survival in peripheral tissues playing a vital role in resistance to infection and cancers. TNFSF18 is found in extracellular space and is integral to membranes and belongs to the TNF ligand superfamily that contains uniform structural motif and the TNF homology domain (THD), which binds to cysteine-rich domains (CRDs) of TNF receptors. TNF can exert many of its effects by binding to cell membrane receptors. The members of the TNF receptor superfamily possess an identical characteristic of an extra cellular domain containing two to six repeats of cysteine rich motifs. The sequence homology between the CRDs and the DNA-binding “zinc-fingers” may be used to speculate intracellular protein phosphorylation by protein kinase C (PKC). IL1R2, also referred to as ILIRB, encodes membrane bound proteins known to be crucially involved in immune response and also functions as a decoy receptor for inflammatory interleukin 1 (IL-1).

Over-expression of ILIR2 has been reported in human uroepithelial cell line (HUC-1) chronically exposed to arsenite, and our results concur with this observation using HaCaT cell line. The cell membrane modulates protein function through localization with the substrate, activator, or downstream target and activation of the protein by a conformational switch. Membrane localized proteins could bind with arsenic at the cell surface and lead to subsequent changes in cellular biological pathways.

The molecular interaction map for ILIB, ILIR2, IL1A, TNFRSF18, and TNFSF18 revealed the E2F4 transcription factor as link (hub) between subnetworks. E2F4 controls cell cycle and function in the suppression of proliferation-associated genes, and its gene mutation and increased expression may be associated with human cancer. E2F4 is predominantly expressed at the onset of keratinocyte differentiation. Further, E2F4 is a promoter of proliferation of human intestinal epithelial crypt cells and colorectal cancer cells. The known predominant expression of E2F4 in the onset of keratinocyte differentiation raises the possibility that arsenic perturbation of protein-protein interactions that include E2F4 could alter keratinocyte differentiation.

From the dataset of 3,829 arsenic-interacting genes from CTD, 106 genes were identified to be enriched for skin pigmentation topics in the ConceptGen enriched biological information (Fig. 3 and Supplementary File 2). Based on our interest in understanding abnormal pigmentation observed in arsenic toxicity, we prioritized two genes, RAC1, a small guanosine triphosphate (GTP)-binding protein, and transferrin receptor (TFRC), an iron uptake receptor, for further discussion.
in this article. According to the Gene Ontology (GO) Cellular Component annotation, \( RAC1 \) and \( TFRC \) are localized to the melanosome.\(^{24} \) \( RAC1 \) was observed in all the data sources (Fig. 2), while \( TFRC \) had been previously identified as possessing vicinal cysteines that are potential binding sites for arsenic.\(^{14} \)

The curated molecular interactions in Comparative Toxicogenomics Database indicate that arsenicals increase \( RAC1 \) activity, localization, or expression (Table 2).

\( RAC1 \) is involved in the signaling process, and actin polymerization is required for the formation of dendrites, which transport the melanosomes from the melanocytic cells to approximately 36 neighboring keratinocytes.\(^{58-60} \) Specifically, \( RAC1 \) mediates formation of lamellipodia (protrusions at the edge of the cell important for cell migration) and melanocyte dendricity.\(^{51,62} \) The transfer of melanosomes to keratinocytes is a mechanism for photoprotection of the skin from ultraviolet (UV) radiation and, thus, the prevention of skin cancer.\(^{63} \) Melanocyte dendricity is critical for skin pigmentation.\(^{64} \) Thus arsenic-induced “increase in localization or activity of the \( RAC1 \) protein” could result in an increase in melanocyte dendricity leading to hyperpigmentation.

The visual analytics–facilitated integration of multiple gene information sources revealed that the transferrin receptor gene (\( TFRC \)) was annotated in the ConceptGen database to be enriched for the melanosome topic. Further, the Comparative Toxicogenomics Database records that in the normal human epidermal keratinocytes (NHEK), sodium arsenite resulted in an increased expression of \( TFRC \) mRNA\(^{13} \) (Fig. 3). We previously showed that \( TFRC \) contain vicinal cysteines using inferences from SNP-induced effects of breakage of disulfide bond.\(^{13} \) Vicinal cysteines in proteins are potential locations for arsenic binding that can affect protein function.\(^{14,65,66} \) Further, \( TFRC \) is a key determinant of the amount and location of iron in the epidermis, and there is a correlation between the expression of \( TFRC \) and iron uptake.\(^{67} \) While 20% to 25% of the absorbed iron is normally eliminated from the body by epidermal desquamation,\(^{68} \) an aberrant \( TFRC \) expression could impair cutaneous iron metabolism leading to the abnormal pigmentation seen in some humans exposed to arsenicals. The environmentally relevant HFE protein associated with hemochromatosis (characterized by excessive dietary iron) resulted in a decreased susceptibility to arsenic trioxide in tumor cell lines. HFE competes with transferrin for binding to the \( TFRC \), which inhibits cellular uptake of transferrin (Fig. 5).\(^{69} \) Reduced iron import (low \( TFRC \) and high HFE) was a signature for favorable prognosis (\( P < 0.005 \)) in breast cancer.\(^{70} \) Changes in iron content of keratinocytes could also be relevant in the susceptibility to arsenic-induced skin cancer.

**Conclusion**

Exposure to inorganic arsenic induces skin cancer and abnormal pigmentation in susceptible humans. High-throughput gene transcription assays such as DNA microarrays allow for the identification of biological pathways affected by arsenic. Investigation of arsenic-induced aberrations in gene expression can help predict initiation and progression of skin cancer and abnormal pigmentation. Molecular network construction for the arsenic upregulated genes \( TNFSF18 \) and \( IL1R2 \) revealed subnetwork interconnections to \( E2F4 \), an oncogenic transcription factor, predominantly expressed at the onset of keratinocyte differentiation. Visual analytics integration of gene information sources helped identify \( RAC1 \), a GTP binding protein, and \( TFRC \), an iron uptake protein as prioritized arsenic-perturbed protein targets for biological processes leading to skin hyperpigmentation. \( RAC1 \) regulates the formation of dendrites that transfer melanin from the melanocytes to neighboring keratinocytes. An increased melanocyte dendricity is correlated with hyperpigmentation. \( TFRC \) is a key determinant of the amount and location of iron in the epidermis. Aberrant \( TFRC \) expression could impair cutaneous iron metabolism leading to abnormal pigmentation seen in some humans exposed to arsenicals. The reported findings contribute to new hypotheses and insights on how arsenic could impair the function of genes and biological pathways related to arsenic toxicity in keratinocytes and melanocytes. Finally, we developed visual analytics resources to facilitate further exploration of the information and the discovery of previously unknown relationships from datasets.

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**Author Contributions**
Conceived and designed the experiments: RDI, UKU, MNA, ANM, ORA. Analysed the data: RDI, UKU, MNA, ANM, MOJ, KF, MAB, RAH. Wrote the first draft of the manuscript: RDI, UKU, MAN, ANM. Contributed to the writing of the manuscript: RDI, UKU, MNA, ANM, MOJ, KF, ORA. Agree with manuscript results and conclusions: RDI, UKU, MNA, ANM, MOJ, KF, MAB, RAH, ORA. Made critical revisions and approved final version: RDI, UKU, MNA, ANM, MOJ, KF, MAB, RAH, ORA. All authors reviewed and approved of the final manuscript.

**Competing Interests**
Authors disclose no potential conflicts of interest.

**Disclosures and Ethics**
As a requirement of publication author(s) have provided to the publisher signed confirmation of compliance with legal and ethical obligations including but not limited to the following: authorship and contributorship, conflicts of interest, privacy and confidentiality and (where applicable) protection of human and animal research subjects. The authors have read and confirmed their agreement with the ICMJE authorship and conflict of interest criteria. The authors have also confirmed that this article is unique and not under consideration or published in any other publication, and that they have permission from rights holders to reproduce any copyrighted material. Any disclosures are made in this section. The external blind peer reviewers report no conflicts of interest.

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