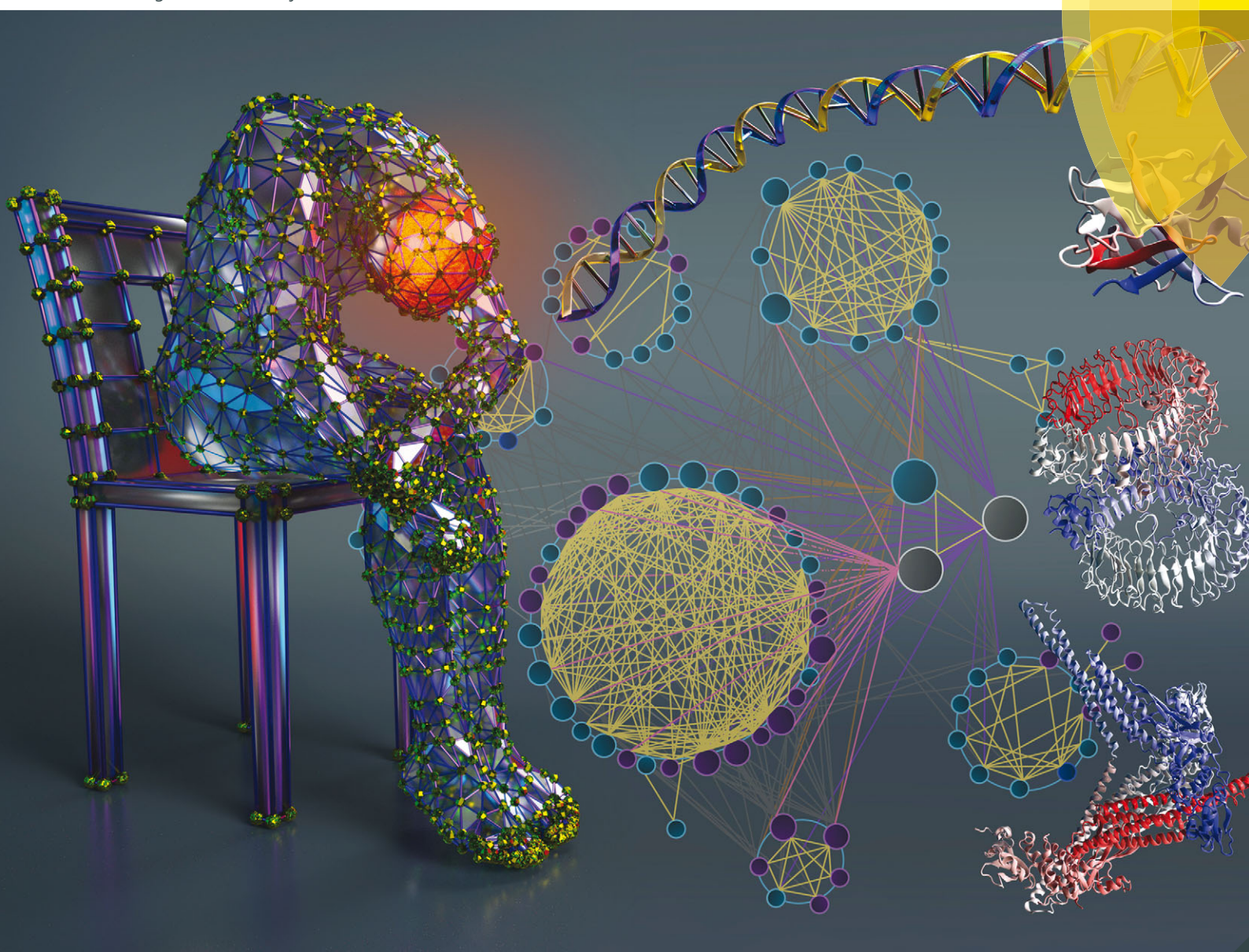


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REVIEW ARTICLE

Francis J. Doyle III *et al.*

Systems biology approach to understanding post-traumatic stress disorder

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www.rsc.org/molecularbiosystemsSystems biology approach to understanding
post-traumatic stress disorderGunjan S. Thakur,^{ab} Bernie J. Daigle Jr,^a Kelsey R. Dean,^{ab} Yuanyang Zhang,^c
Maria Rodriguez-Fernandez,^{ab} Rasha Hammamieh,^d Ruoting Yang,^d Marti Jett,^d
Joseph Palma,^{ad} Linda R. Petzold^{be} and Francis J. Doyle III^{*ab}

Post-traumatic stress disorder (PTSD) is a psychological disorder affecting individuals that have experienced life-changing traumatic events. The symptoms of PTSD experienced by these subjects—including acute anxiety, flashbacks, and hyper-arousal—disrupt their normal functioning. Although PTSD is still categorized as a psychological disorder, recent years have witnessed a multi-directional research effort attempting to understand the biomolecular origins of the disorder. This review begins by providing a brief overview of the known biological underpinnings of the disorder resulting from studies using structural and functional neuroimaging, endocrinology, and genetic and epigenetic assays. Next, we discuss the systems biology approach, which is often used to gain mechanistic insights from the wealth of available high-throughput experimental data. Finally, we provide an overview of the current computational tools used to decipher the heterogeneous types of molecular data collected in the study of PTSD.

^a Institute of Collaborative Biotechnologies, University of California Santa Barbara,
Santa Barbara, California, USA^b Department of Chemical Engineering, University of California Santa Barbara,
Santa Barbara, California, USA. E-mail: doyle@engineering.ucsb.edu;
Fax: +1-805-893-4731; Tel: +1-805-893-8133^c Department of Computer Science, University of California Santa Barbara,
Santa Barbara, California, USA^d US Army Center for Environmental Health Research, Fort Detrick, Maryland, USA^e Department of Mechanical Engineering, University of California Santa Barbara,
Santa Barbara, California, USA

1 Introduction

Post-traumatic stress disorder (PTSD) is the fifth most common psychiatric disorder with an occurrence rate of about 8% in the United States.¹ Unlike other psychiatric disorders, PTSD may develop in individuals only after they have experienced a terrifying traumatic event such as torture, rape, kidnapping, or natural disasters such as floods or earthquakes. The incidence of



Gunjan S. Thakur

interests include dynamical systems, system biology, controls, machine learning and mathematical modeling of biological systems.

Gunjan Singh Thakur is a post-doctoral scholar in the Institute for Collaborative Biotechnologies at University of California Santa Barbara. He received his Bachelor of Technology from Indian Institute of Technology, Guwahati in Mechanical Engineering in 2000, Masters of Science in Engineering Science and Mechanics from Virginia Polytechnic Institute and State University, in 2004 and PhD in Mechanical Engineering from UCSB in 2011. His research



Bernie J. Daigle Jr

the recipient of Geraldine Jackson Fuhrman and Howard Hughes Medical Institute Predoctoral Fellowships. His research interests include computational modeling and analysis of stochastic biochemical systems, and integration of genome-scale datasets for disease biomarker identification.

Bernie Daigle, Jr is Assistant Researcher in the Institute for Collaborative Biotechnologies at the University of California, Santa Barbara. Prior to his current appointment, he was a postdoctoral researcher in the Department of Computer Science at UCSB. He received his BS in Biology, summa cum laude, from Cornell University in 2002. In 2010, he received his PhD in Genetics from Stanford University School of Medicine, where he was

occurrence of PTSD in war veterans is relatively high. Approximately 22% of veterans returning from Iraq and Afghanistan are diagnosed with the disorder.² Before 1980, when it was officially recognized as a medical disorder, PTSD was referred to by different names. During the American Civil War it was referred to as “Soldier’s Heart”, during World War I as “Combat Fatigue”, during World War II as “Gross Stress Reaction”, and during the Vietnam War as “Post-Vietnam Syndrome”. Other names for PTSD include “Battle Fatigue” and “Shell Shock”.

Currently, survey-based assays are used to diagnose PTSD, including the clinician-administered PTSD scale (CAPS)³ from

the “Diagnostic and Statistical Manual of Mental Disorder” (considered to be the gold standard) and self assessment scales such as the “Posttraumatic Diagnostic Scale”.⁴ These tests help diagnose whether or not a subject has PTSD with fairly good accuracy. However, these tests are not designed to diagnose stages of the disorder, from an early manifestation to later advanced stages. Thus there is a need to identify reliable biomarkers for PTSD that can be used for the accurate diagnosis, prognosis, and evaluation of therapeutics for the disorder. Although many candidate biomarkers have been identified for PTSD, reliable markers for specific clinical applications are still lacking.



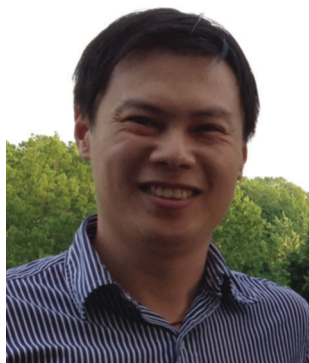
Kelsey R. Dean

Kelsey R. Dean is a third year chemical engineering graduate student at the University of California Santa Barbara, where she was awarded the Heslin Fellowship in 2012. She received her BS in Chemical and Biomolecular Engineering from Johns Hopkins University in 2012. Her research interests include unsupervised learning techniques for disease subtype identification, and clinical and biological data integration for disease biomarker identification.



Rasha Hammamieh

Dr Hammamieh is the Director of Integrative Systems Biology at US Army Center for Environmental Health Research. Her work focuses on the development of diagnostic biomarkers and therapeutic strategies for diseases of military relevance, including PTSD, exposure to pathogens, battlefield-like stress, coagulopathy and pain. Dr Hammamieh has authored and co-authored over 40 publications on responses to stressors including pathogens, trauma, PTSD, battlefield-like stress and nutrigenomics.



Ruoting Yang

Ruoting Yang is Bioinformatics Supervisor in the Frederick National Laboratory for Cancer Research. Prior to his current appointment, he was a post-doctoral researcher in the Institute for Collaborative Biotechnologies, University of California Santa Barbara. He received his BS in Automation, from Xiamen University in 2000, and MS in Control Theory in Academy of Mathematics and Systems Science, Chinese Academy of Sciences. In

2008, he received his PhD in Systems Science and Mathematics and MS in Electrical Engineering from Washington University in Saint Louis. His research interests include systems biology, computational biology, mathematical modeling, and biomarker identification.



Francis J. Doyle III

Dr Doyle is the Chair of the Department of Chemical Engineering at UCSB, and is Director of the Army Institute for Collaborative Biotechnologies. He holds the Duncan and Suzanne Mellichamp Chair in Process Control in the Department of Chemical Engineering, as well as appointments in the Electrical Engineering Department, and the Biomolecular Science and Engineering Program. He received his BSE from Princeton (1985), C.P.G.S. from Cambridge (1986), and PhD from Caltech (1991), all in Chemical Engineering. He is the recipient of numerous research awards (including the NSF National Young Investigator, ONR Young Investigator, Humboldt Research Fellowship) as well as teaching awards (including the Purdue Potter Award, the ASEE Ray Fahien Award, and the ASEE Chemstations Lectureship Award). He is a Fellow of multiple professional societies including IEEE, IFAC, AIMBE, and AAAS. His research interests are in systems biology, network science, modeling and analysis of circadian rhythms, drug delivery for diabetes, and model-based control.

In a number of disease models, system biology approaches have shed light on the complex underlying biophysical networks.^{5–8} Systems biology aims to understand the complexity of a biological system by integrating experimental data at different scales with mathematical models (*e.g.*, differential equations, stochastic models and data-driven approaches such as machine learning) to obtain useful information about the system at the scale of interest.⁹ These scales range from the single-cell and single-molecule level up to the genome-wide, organismal level (see Fig. 1). Depending on the scale of investigation, informative experimental data may consist of “omics” measurements (*e.g.*, transcriptomics and proteomics), quantification of individual metabolites, or brain images. The systems approach enables us to make testable predictions (*e.g.*, identification of diagnostic/prognostic biomarkers or drug targets) and gain mechanistic insights into the system in question.

In the remainder of this review, we first provide a brief overview of the known biological underpinnings of PTSD, noting that many excellent reviews are available reflecting the vast, but often conflicting, amount of information available on the subject.^{10–14} Next, we focus on the range of systems biology approaches that have been used to understand the disorder. Finally, we present model studies which have employed these tools to obtain new clinical and biological insights.

2 Biology of PTSD

A large body of literature has documented symptoms of PTSD in human subjects. These symptoms include memories/avoidance of traumatic events, emotional numbing, hyper-arousal

(*e.g.* exaggerated startle and lack of sleep), alteration in fear conditioning, and impaired extinction learning.^{15–17} Recently, many neurobiological studies have been performed, effectively broadening the view that PTSD is purely a psychological disorder and helping to obtain a deeper insight into the disease mechanism. Psychophysiological observables such as heart rate,¹⁸ skin conductance (measurement of sweat activity), eye blinking (measurement of startle state), facial electromyogram (EMG, used to assess the neuronal signaling to muscles) and cortical electroencephalographic event related potentials (ERPs, measurement of brain activity)^{19,20} have been used to quantify PTSD symptoms in a systematic manner. Most psychophysiological studies conduct cross-sectional comparisons between the treatment (individuals with PTSD) and control (individuals without PTSD) groups. These studies generally find autonomic reactivity, such as heart rate and skin conductance, and startle response to trauma-related stimuli to be elevated in the treatment group. Specifically, a strong correlation has been observed between the level of reactivity to trauma-related stimuli and the severity of PTSD.^{21,22} Increases in heart rate, recurrent recall of trauma memories, and exaggerated startling are typically all acquired in individuals with the disorder, but these behaviors may not be unique to PTSD.

Neuroimaging

To gain a better understanding of the disorder, many studies have focused on macroscopic features of the human brain by employing techniques such as structural (sMRI) and functional magnetic resonance imaging (fMRI). The brain regions related to symptoms of the disorder include the hippocampus, amygdala, ventromedial prefrontal cortex (vmPFC), dorsal anterior cingulate

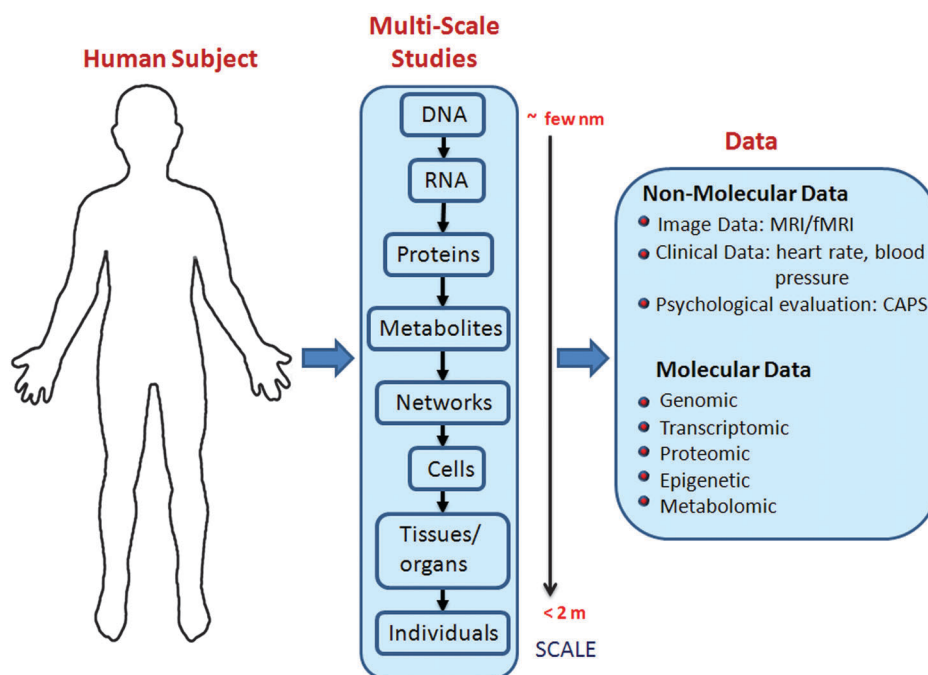


Fig. 1 Systems biology approach for the study of PTSD: a multitude of data types can be experimentally collected by probing biological systems at different scales. This wealth of data is then further processed by employing mathematical and computational techniques (*e.g.*, statistics, machine learning, computational modeling) to gain insight into the system under study.

Table 1 Summary of neuroimaging findings in subjects with PTSD. Abbreviations used: vmPFC denotes ventromedial prefrontal cortex and dACC denotes dorsal anterior cingulate cortex

| Brain regions | Function imaging | | | Structural imaging |
|----------------|--|-------------------------------|--------------------------------|---------------------------|
| | Normal function | Activity in PTSD subjects | Correlation with PTSD severity | Volume change |
| Amygdala | Detection of threat, processing/learning of fear stimulation ^{47,48} | Increased ^{32,33} | Positively ^{44–46} | Not clear |
| vmPFC | Goal-directed decisions making ⁴⁹ | Decreased ^{36,44,50} | Negatively ^{44,51} | Decrease ²⁵ |
| dACC | Regulates cognitive control, error detection, fear appraisal/expression ^{54,55} | Increased ^{51–53} | Positively ^{34,38} | Decrease ^{25,26} |
| Hippocampus | Short and long term memory ^{56,57} | Mixed ^{58,59} | Negatively ²⁹ | Decrease ^{23,24} |
| Insular cortex | Monitoring internal body states ⁶⁰ | Increased ^{35,61} | Positively ³⁵ | Decrease ³⁵ |

cortex (dACC) and insular cortex (see Table 1). sMRI studies have shown a decrease in the volume of hippocampus,^{23,24} vmPFC,²⁵ dACC^{25,26} and the insular cortex^{27,28} for patients with PTSD. In the case of changes to the hippocampal volume, it is not clear whether this decrease is caused by the extreme trauma or simply represents a risk factor; however, some studies point to the latter.²⁹ We note that exceptions to the above have been reported in children with PTSD,³⁰ possibly due to lack of neuronal maturation, and in cases where symptom severity was not very strong.³¹ Studies performed with combat twins using voxel-based morphometry suggest that the reduction in ACC volume may be an acquired feature due to extreme trauma.²⁵

Functional MRI studies have shown increased activity in the amygdala,^{32,33} dACC³⁴ and insular cortex,³⁵ and decreased activity in the vmPFC³⁶ for PTSD patients subjected to trauma-related stimuli. Positron emission tomography, an fMRI technique quantifying local changes in cerebral blood flow, has been performed on identical twins to study resting dACC glucose metabolism and dACC activation during non-emotional inference tasks. These studies demonstrated that combat-exposed veterans with PTSD and their co-twins have greater metabolism³⁷ and greater activation³⁸ compared to veterans without PTSD and their co-twins. This suggests that increased activation of dACC is a PTSD risk factor rather than a symptom. Finally, activation in the amygdala³⁹ and insular cortex⁴⁰ does not appear to be unique to PTSD, as these responses are also seen in other anxiety disorders. Functional MRI techniques are also used to measure resting state brain activity and functional connectivity (*i.e.*, temporal correlations between brain regions with common functional properties). A recent paper by Yan *et al.*⁴¹ found increased activity in the amygdala and the anterior insula and decreased activity in the thalamus to be common features in both resting state brain activity and task-based fMRI studies (as reported by other authors, see citations included in the paper), in male combat veterans with PTSD. Their results also highlight decreased activity in the precuneus region of the brain, which is responsible for integrating information from the past and future. Furthermore, Rabinak *et al.*⁴² found enhanced connectivity between the amygdala and insula in combat-related PTSD patients. However, as many studies of anxiety-related disorders have also reported fMRI-based amygdala and insula activation,⁴³ these responses may not be unique to PTSD.

Neuroendocrinological studies

The major constituents of the neuroendocrine response to physical and emotional threat and stress are the sympathetic nervous system (SNS) and the hypothalamic-pituitary adrenal (HPA) axis.⁶² The immediate SNS response mobilizes the acute fight-or-flight response and is followed by a response from the HPA axis that reinstates homeostasis. PTSD patients showed low cortisol levels, which is surprising for a disorder precipitated by extreme stress, and differs from that observed in studies of acute and chronic stress and major depressive disorder (which have been associated with increased cortisol levels).⁶³ Moreover, PTSD patients exhibited increased levels of norepinephrine, corticotropin-releasing hormone (CRH) and proinflammatory cytokines, reflecting reduced glucocorticoid signaling. This profile of alterations has been associated with PTSD pathophysiology. However, emerging research indicates that these alterations may instead reflect pre-traumatic vulnerabilities to the later development of PTSD.⁶²

Gene expression

Previous gene expression studies of PTSD have focused on two types of biomarkers: predictive and diagnostic signatures. Predictive biomarkers indicate risk of PTSD development after trauma exposure and require longitudinal studies. Multiple longitudinal studies have presented panels of potential risk biomarkers.^{64,65} These biomarkers include up-regulated genes involved in type-1 interferon signaling (IFI27, OAS1, OAS2, OAS3, XAF1, and USP18)⁶⁴ and differentially expressed genes which associate with glucocorticoid receptors (GR), including low FKBP5 and high GILZ expression.⁶⁵ Additionally, many studies suggest potential mRNA diagnostic biomarkers, including the GR-associated genes FKBP5 and STAT5B⁶⁶ among others.^{67,68} Early markers for PTSD development have been reported which may aid in developing early detection diagnostic tools. Gene expression data collected in emergency rooms immediately following trauma led to the discovery of 574 differentially expressed genes distinguishing those who would later develop PTSD from those who would recover.⁶⁹ These transcripts were enriched in genes involved in immune activation, cell cycle, and signal transduction, among other biological processes. While these and other initial results are promising, no validation studies have shown that these biomarkers are valid in

independent data sets. Additionally, the biomarkers were identified strictly by differential expression statistics or panels derived from classification algorithms (e.g., support vector machines) and do not incorporate known or inferred biological network information. This limits the ability of these gene panels to give biological insight into the development and progression of PTSD and does not provide a clear path for identification of therapeutics.

Epigenetics

Research on epigenetic mechanisms has revealed that environmental experiences such as early life stress and fear conditioning can modify the expression of genes without altering the corresponding DNA sequence.^{11,70–73} Most of the studies investigating the role of epigenetics in PTSD have focused on DNA methylation, typically comparing peripheral blood methylation levels between PTSD patients and controls.^{74–78} In particular, studies have found that the number of uniquely methylated genes was significantly higher in PTSD-affected patients.⁷⁹ Furthermore, genes found to be hyper-methylated in PTSD patients include: COMT,⁸⁰ H19,⁸¹ IL18,⁸¹ LINE-1,⁸² PAC1 (female),⁸³ SLC6A3,⁸⁴ MAN2C1,⁸⁵ CLEC9A,⁸⁶ ACP5,⁸⁶ and TLR8.⁸⁶ Conversely, genes such as FKBP5,⁸⁷ Alu,⁸² SLC6A4,⁷⁴ TPR, and ANXA2⁸⁶ were found to be hypo-methylated in PTSD. Studies have also found that for subjects with low socioeconomic status, the relationship between DNA methylation and risk of developing PTSD may be modified in genes predominantly related to nervous system function.⁸⁸

Subtypes of PTSD

The heterogeneous nature of the symptoms and development of PTSD create challenges in diagnosing and treating the disease. While multiple studies have emphasized clear divisions into PTSD subgroups,^{89,90} very little work has focused on identifying unique biological signatures for these subtypes. Instead, recent work has focused on using symptoms and symptom trajectories to identify these distinct subgroups, or on correlating specific biological measures to these symptom classes. For example, longitudinal data from the Jerusalem Trauma Outreach and Prevention Study identified three trajectories of PTSD symptoms which they called rapid-remitting, slow-remitting and non-remitting.⁸⁹ Interestingly, they note that only subjects classified as slow-remitting showed symptom improvements when given antidepressants. Similar responses to antidepressants were seen in another study⁹⁰ where promoter methylation of FKBP5 and GR showed differences between responders and non-responders to treatment. These examples further emphasize the existence of subgroups within PTSD, but overall provide little biological insight into the key differences between these groups. In order to progress in the identification and characterization of these subtypes in a manner useful for diagnostic and therapeutic development, specific biological signatures must be extracted from large-scale data. In particular, the biological features extracted from this type of analysis would provide a more comprehensive picture of subtype biology than these single gene correlations. We note that the identification of

subgroups from DNA microarray data has previously been applied to the study of diseases such as cancer and fibromyalgia.^{91–93}

3 Systems biology approach

The molecular inputs used for understanding a given biological system from a systems perspective include genomic, transcriptomic, epigenomic, metabolomic, and proteomic data. These data types measure the genome-wide amounts of and changes to DNA sequence, mRNA expression, DNA methylation/histone modification, metabolite production, and protein expression, respectively. Commonly used experimental techniques for collecting this data include high-throughput sequencing methods, array-based hybridization assays (e.g. microarrays), liquid and gas chromatography, and mass spectrometry. In addition to high-throughput molecular measurements, neuroimaging assays, such as functional and structural MRI, provide an organism-wide measure of brain activity and morphology, respectively. When studying disease mechanisms, each of the above assays can be utilized in either a cross-sectional or longitudinal manner. In the former case, different subjects with and without the disease phenotype are measured simultaneously, with the goal of identifying time point-specific differences between the groups. In the latter case, measurements are repeatedly collected from the same subjects over time, which helps to reduce inter-individual variability and can provide temporal information regarding disease progression. Fig. 2 shows the analysis pipeline typically applied to “omics” data to uncover the underlying biology of PTSD. In the remainder of this section, we review some commonly used methods in this pipeline.

Raw data from all of the above experimental modalities are corrupted with noise. It is essential to understand the sources of this noise in order to control for them in downstream analyses, as the final conclusions of the study depends on it. Noise sources can be broadly categorized into two groups: one deriving from the stochastic nature of molecular interactions (referred to as biological noise), and the second coming from the measurement techniques and other experimental strategies used (measurement noise). As an example of the first group, gene expression levels measured in genetically identical cells show substantial cell-to-cell variations.^{94,95} These unavoidable variations are attributed to random molecular interactions and are known to influence phenotypic properties,⁹⁶ environmental response,⁹⁷ decision making at different stages of cell development,⁹⁸ and information processing.⁹⁹ In contrast, the measurement noise introduced into the data must be minimized, otherwise it can lead to incorrect downstream analyses.¹⁰⁰

As the first step of data analysis, the raw data is corrected for measurement noise. Methods for performing this task with high-throughput data include background correction, normalization and batch correction. The first technique is typically used in fluorescence-based assays for reducing the effects of non-specific background fluorescence. One part of this procedure often involves removing those measurements that are indistinguishable from background noise. Although this introduces missing

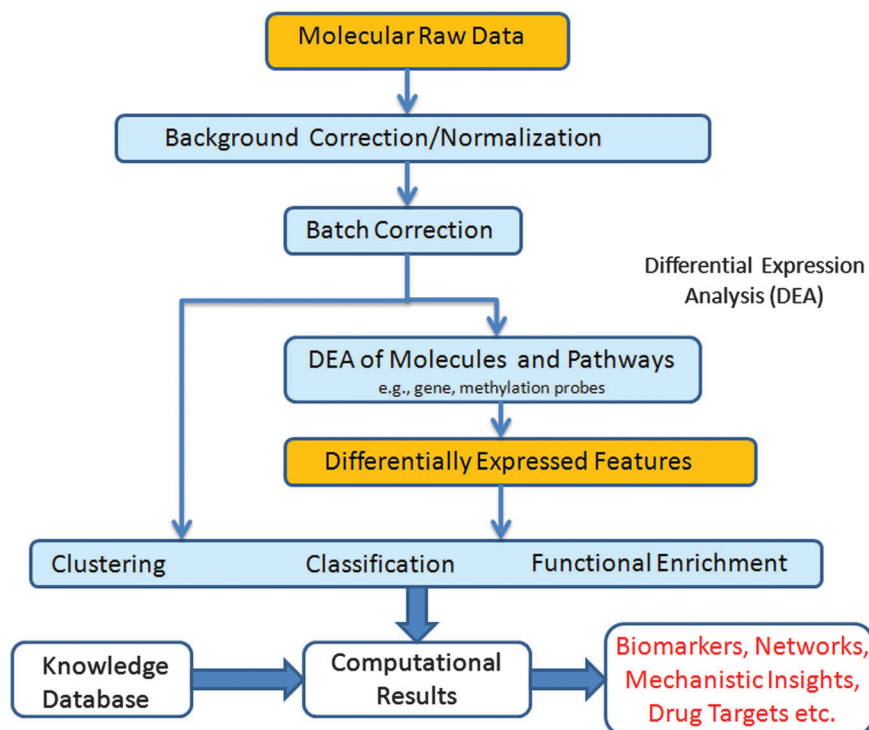


Fig. 2 Analysis pipeline for heterogeneous molecular data generated in PTSD studies. The raw data is first corrected for systematic errors and batch effects, both of which may be introduced during the experimental process. Provided the biological signal is significant, correcting these errors improves the signal-to-noise ratio and ensures more reliable downstream analyses. Following this step, statistical and machine learning tools are used at appropriate stages (shown in the blue boxes) to obtain useful system information (shown in red).

values into the dataset, such values are preferable to measurements entirely lacking in biological signal. The second technique, normalization, is used in most high-throughput assays to remove systematic biases in the data due to experimental artifacts rather than biological signal. Examples of such biases include differing amounts of biological starting material across assays (between-assay normalization) and unequal fluorescence labeling efficiencies in dual-color fluorescence-based assays (within-assay normalization). Normalization methods often make assumptions about the biological signal present in the data, including that the overall signal range is the same across all samples and that the majority of assayed measurements do not differ significantly between samples.¹⁰¹

Variability in high-throughput data is often associated with location,¹⁰² time,¹⁰³ individuals conducting the experiment and other technical factors.^{104,105} In addition to measurement noise, this gives rise to what is known as a batch effect—where the structure of measurement error, generally thought to be fixed within each batch, varies across batches. This effect must be taken into account (see the second step in Fig. 2) to allow the integration of data from different batches, thus obtaining greater statistical power for further analyses.¹⁰⁶ Methods such as surrogate variable analysis¹⁰⁷ (SVA), independent surrogate variable analysis¹⁰⁸ (ISVA), and ComBat¹⁰⁶ have been used to perform batch correction. ComBat is an empirical Bayes method which is robust to outliers and can also correct for small batch size. When the sources of experimental batches are simple¹⁰⁰

(e.g., processing time), ComBat outperforms many other methods.¹⁰⁹ However, certain confounding factors are more difficult to ascertain. For these cases, SVA and ISVA provide methods to correct for noise due to unmodelled, unmeasured, or unknown factors. However, we note that these methods do not address the situation where the phenotype of interest is highly correlated with the experimental batches.

Once noise correction has been performed on the available high-throughput data, a range of analysis methods can be applied to characterize the biological signal present. One of the most common methods used to study disease mechanism is differential expression (DE) analysis. The goal of DE analysis in this context is to identify genes/metabolites/proteins that are differentially expressed between disease and control samples. Use of proper statistical testing is essential for this task, as the huge numbers of candidate biomarkers in genome wide studies (e.g., roughly 20 000 genes on a typical microarray) can lead to large numbers of false positive and false negative predictions. Statistical methods like the classical *t*-test, moderated *t*-test¹¹⁰ and SAM statistic¹¹¹ have all been successfully used to generate DE *p*-values (i.e., probabilities that candidate biomarkers' DE scores are due to random chance rather than biological signal). Given a list of *p*-values, multiple hypothesis testing correction methods that control the false discovery rate (FDR) or family wise error rate (FWER) can then be used to reduce numbers of false positives and negatives.¹¹² After identifying disease-specific DE candidates, a natural next step is to perform gene

set enrichment analysis (GSEA).¹¹³ Using a Kolmogorov–Smirnov (K–S) statistic, GSEA identifies statistically enriched biological functions or pathways within a list of candidate biomarkers. GSEA can be very useful for ascribing biological meaning to output from DE analyses.

An alternative to DE analysis for identifying disease biomarkers is to employ machine learning tools. In general, there are two distinct algorithmic approaches, namely (1) supervised and (2) unsupervised learning. The nature of the available data and the objective of the learning task dictates the selection of the proper approach. Unsupervised learning is used when observation labels (*e.g.*, class or category) are not available. Algorithms of this type are widely used for clustering, dimensional reduction, principal component analysis, and probability density estimation of a population. In contrast, supervised learning refers to algorithms that use available observation labels in the learning process. These algorithms are mostly used for classification (*i.e.*, assigning membership labels to a given observation) and regression (*i.e.*, estimating relationships between variables). Supervised classification generally consists of two phases: training and testing. During the training phase, features from samples of known status (*e.g.*, disease or control) are used to build a predictor that is then evaluated during the testing phase on samples of unknown status. Many algorithms are available for classification, including: decision trees, random forests, k-nearest neighbors, Bayesian methods, neural networks, support vector machine (SVM), nearest shrunken centroids (NSC),¹¹⁴ pathway-based classifiers like the condition-responsive genes (CORGs) method,¹¹⁵ and core module biomarker identification with network exploration (COMBINER).¹¹⁶ However, only a limited number of these algorithms have been previously used for PTSD research (*e.g.*, SVM,⁶⁴ Logistic regression,⁶⁵ COMBINER¹¹⁶). Currently used methods for PTSD identify biomarker panels by considering combinations of single, unrelated genes or gene sets associated with known pathways. However, none of these methods utilize known molecular interaction networks, which can provide useful insight into network perturbations associated with PTSD, and has shown improvement in classification performance and stability for diseases such as cancer.¹¹⁷ Generally, classification performance can be quantified in several ways; some of the most common include: prediction error rate, sensitivity, precision, F-score and area under the receiver operating characteristic curve (AUC). The choice of which specific classification algorithm to use is critical, as an incorrectly chosen classifier often leads to poor performance. In this case, one must try to select a more appropriate algorithm given factors such as size of the training set, preselection of features given as input to the algorithm, the dimensionality of the dataset, and a potential imbalance in class sample sizes. In addition, algorithm parameters can often be more optimally tuned to improve classification performance. Further information addressing the above can be found in published reviews.^{118–121}

In order to derive mechanistic insights for a given condition, the systems biology approach also attempts to predict interaction networks between different chemical species. To do this, the expression levels of genes, proteins, and metabolites are

first measured using high-throughput technologies. Depending on the nature of the species under study, different types of interaction networks can be inferred. For example, network analysis of gene expression data gives rise to gene regulatory networks, while a similar analysis of protein expression data generates signal transduction networks. The inferred relationships between species range from physical interactions (*e.g.*, between transcription factors and their target genes) to functional interactions (*e.g.*, correlated expression between groups of genes). Statistical methods such as regression and correlation analysis,¹²² mutual information-based approaches,¹²³ Bayesian networks,¹²⁴ and probabilistic graphical models¹²⁵ have all been used to predict interactions between species. Detailed reviews of this topic can be found in the following ref. 122, 124, 126–129.

An important challenge in the systems biology approach is the fusion of information from multiple “omics” sources.¹³⁰ Zhao *et al.* have used machine learning techniques to perform unsupervised gene biomarker selection in cancer by integrating both mRNA and miRNA data.^{131,132} Ye *et al.* have identified Alzheimer's disease biomarkers by combining MRI data, demographic information, and genetic information.¹³³ Finally, Xiang *et al.* have identified Alzheimer's biomarkers by combining proteomics, PET, MRI and CSF datasets.¹³⁴ However, to our knowledge there are no studies performing heterogeneous data integration for the analysis of PTSD.

4 Model studies

In this section we review representative studies employing systems tools to investigate PTSD. The first two studies use ordinary differential equation (ODE) models to obtain a mathematical description of the disorder, while those remaining use statistical and machine learning techniques.

Subjects with PTSD are known to have altered fear conditioning responses. As a first case study we highlight the work done by Li *et al.*,¹³⁶ where they performed systems modeling of neurons in the lateral amygdala (LA) region (a key site of synaptic events that contribute to fear learning). Using experimental data from the literature, the authors built a computational network model consisting of ten neurons found in the amygdala—eight pyramidal cells and two GABAergic interneurons. Their twin-compartment model represents a soma (diameter of 15 μm ; length of 15 μm) and the dominant apical dendrite. The authors derived ODEs for somatic and dendritic membrane potentials by balancing different current sources (*e.g.* intrinsic, synaptic currents and electrode current). They then used their model to demonstrate the characteristic abilities of the LA in both fear conditioning and extinction. Their results suggest that fear expression is determined by a balance between pyramidal cell and interneuron excitations. Furthermore, the authors show that fear memory is stored in the pyramidal cells, whereas extinction memory is stored in both interneurons and pyramidal cells. Finally, results from the model suggests that fear memory is not erased fully by extinction; rather, it is inhibited by interneurons

that undergo synaptic plasticity during extinction training. As emphasized by the authors, their model for the plasticity of inhibitory synapses in the LA improves understanding of the fear mechanism.

Mathematical models of varying complexity have been proposed to describe the circadian dynamics of the HPA axis (see Vinther *et al.*¹³⁷ and references therein). An ordinary differential equation model for cortisol dynamics in the HPA axis has been developed by Sriram *et al.*¹³⁵ (see Fig. 3), where they seek to understand the mechanism leading to hypocortisolemia in PTSD and hypercortisolemia in depressed patients. The developed model and its estimated parameters support Yehuda's hypothesis that hypocortisolemia in PTSD is due to the strong negative feedback loop operating in the neuroendocrine axis under severe stress.¹³⁸ The model was used to simulate *in silico* patients whose behavior suggested that, due to disrupted negative feedback loops, cortisol levels are different in normal, PTSD and depressed subjects during the night. Moreover, the model predicted transitions from normal to diseased states due to changes in the strength of the negative feedback loop and the stress intensity in the neuro-endocrine axis.

Animal models have been used to investigate the effects of trauma and draw parallels with PTSD and other neurodegenerative disorders in humans. Next we consider a recent study by Hammamieh *et al.* where they developed a "social-defeat" model by exposing male C57BL/6J mice to aggressor mice for six hours daily for five or 10 days in total, using a "cage-within-cage resident-intruder" protocol.¹⁴⁰ The mice showed significant behavioral, physiological, and histological changes associated with the trauma stressor. Furthermore, seven brain regions, including the hippocampus, amygdala, and medial prefrontal cortex, as well as whole blood were

harvested and the corresponding whole-cell mRNA populations were extracted. In order to understand the underlying molecular principles, Yang *et al.*¹³⁹ used a computational tool entitled core module biomarker identification with network exploration (COMBINER). This approach using machine learning and statistical tools, was first developed by Yang *et al.*¹¹⁶ COMBINER provides a novel method for the identification of disease biomarkers. Designed to take multiple data cohorts as input, COMBINER identifies gene module biomarkers that are consistent across the population of samples and constructs their associated regulatory networks. The social defeat study identified common differentially expressed genes in multiple brain regions as well as between the brain and blood. These DEGs were used to obtain the core disease module (see Fig. 4). Many of the resulting modular gene networks were found to be previously associated with PTSD, depression, and other neurodegenerative diseases.

Studies^{141,142} have indicated that subjects with PTSD also have a higher risk of cardiovascular conditions, including increased basal heart rate and blood pressure, higher risk for hypertension and stroke. A recent study by Cho *et al.*¹⁴³ used a systems approach to study the effect of traumatic experiences on heart function using the animal model developed by Hammamieh *et al.*¹⁴⁰ In particular, they found that myocarditis was a frequently detected pathology in stressed subject mice, suggesting a stress-associated heart injury. The authors studied the temporal expression of mRNA and miRNA in heart samples obtained from stressed and control animals, providing a characterization of the heart tissue repair process in response to stress-induced tissue injury. Inflammatory-related genes were found to be significantly regulated after as little as one day of exposure to stress, and most of the changes in gene expression returned to normal levels after a recovery period. Additionally, key molecular processes involved in classical wound healing in the heart tissues of subject mice were identified, including the chemokine signaling pathway, extracellular remodeling and epithelial to mesenchymal transition (EMT). Taken together, the results of Cho *et al.*¹⁴³ suggest a molecular basis for the observed higher risk of cardiovascular disorders in PTSD patients, further illustrating the increased likelihood of cardiac dysfunction induced by long-term stress exposure.

Blood samples are one of the most common sources for obtaining biological data. A gene expression study of the peripheral blood mononuclear cells (PBMC) has been performed by Segman *et al.*⁶⁹ In this study, blood samples were collected for subjects visiting the emergency room following trauma as well as after a four month follow-up visit. The aim of the study was to identify the PTSD gene expression signature by applying differential expression analysis, classification, and functional enrichment of the blood samples collected. Differential expression analysis identified DEGs as genes which were significantly ($p < 0.05$) expressed using three methods: *t*-test, threshold number of misclassifications (TNOM), and the Info¹⁴⁴ method. The number of DEGs identified by the intersection of these three methods is 408, 574, and 656 genes for the emergency room, four month follow-up, and combined times

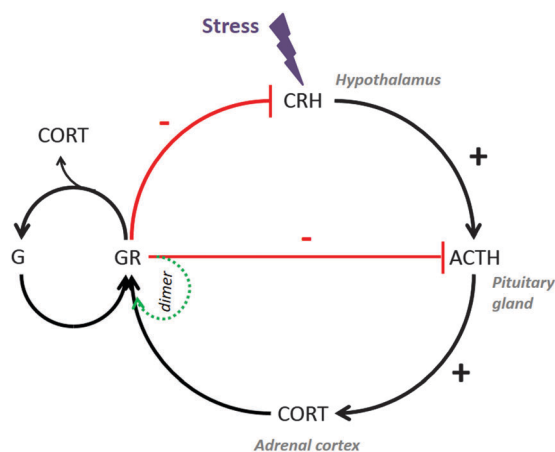


Fig. 3 Regulatory network of cortisol in the HPA axis. Stress induces the secretion of corticotropin release hormone (CRH) in the hypothalamus that in turn activates the release of adreno-corticotropin hormone (ACTH) in the anterior pituitary gland. ACTH moves to the adrenal cortex and stimulates cortisol (CORT). The secreted cortisol binds to glucocorticoid receptors (G) to form a complex GR followed by dimerization. Cortisol complex GR binds to both CRH and ACTH to down regulate the production of cortisol. This closed loop gives rise to a negative feedback in the circuit that is vital in maintaining the homeostasis of the system during stress.¹³⁵

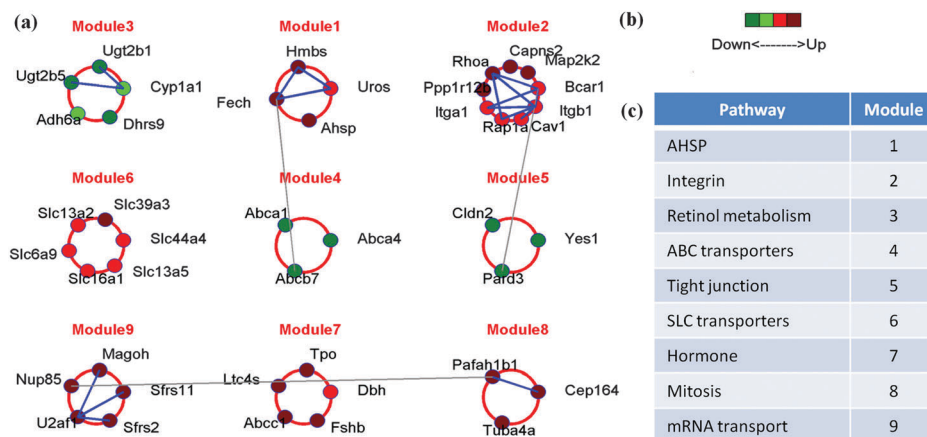


Fig. 4 Blood-brain gene network for a “social defeat” mouse model. (a) Nine core gene expression modules identified by COMBINER. The core modules consist of only those genes that are expressed in both blood and multiple brain regions (which include the amygdala, hippocampus, hemibrain, medial PFC, septal region, stria terminalis and ventral striatum). Known protein–protein interactions (PPIs) are marked by lines connecting genes—blue lines denote within-module interactions, while gray lines denote between-module interactions. (b) The color scale for the blood expression level of each gene in the nine modules, as indicated by the colored circles in (a). (c) Putative biological functions of the expression modules as inferred using KEGG annotation. The above figure has been adapted from Yang *et al.*¹³⁹

points, respectively. Next, a Naive Bayesian Classifier was trained and tested using leave-one-out cross validation (LOOCV) on the samples from the emergency room (ER) (6 PTSD, 5 control) and the four month follow-up (5 PTSD, 4 control). The ER and follow-up classifiers had misclassification error rates of 18% and 11%, respectively with all misclassifications being false positives. Additionally, gene ontology enrichment analysis was used to understand the biological pathways related to the identified PTSD signature. Terms significantly enriched in DEGs from the differential expression analysis were determined using the hypergeometric *p*-value and a false discovery rate (FDR) cutoff of 0.1. Terms related to RNA processing, metabolism, and binding as well as nucleic acid metabolism and binding were found to be significant. Overall, this exploratory analysis presented both gene and pathway-level analysis of PTSD mRNA signatures from blood, the result of which require further validation in larger studies to understand their usefulness as molecular markers of PTSD.

In addition to gene expression signatures in blood, there has been increasing interest in epigenetic markers of PTSD, specifically DNA methylation. The changes in DNA methylation can occur during development, but also later in life in response to an individual's environment and experience. These epigenetic changes can create long-term changes in gene expression, resulting in disease phenotypes involving perturbed biological function. The first published epigenetic study of PTSD⁷⁹ identified genes which were uniquely methylated or unmethylated in those who developed PTSD, compared to control subjects who had experienced similar levels of trauma. This exploratory study of methylation in 14 000 genes over 23 PTSD and 77 control samples identified approximately 400 genes which were uniquely methylated or unmethylated in PTSD subjects using the Wilcoxon test ($p < 0.01$). To understand the related biology, the authors used functional annotation clustering (FAC) analysis to identify Gene Ontology terms that were enriched in the set of

uniquely methylated and unmethylated genes. Terms related to both the innate and adaptive immune system were identified as the most significantly enriched. This finding was further strengthened by an independent measure of immune function: the amount of antibodies to a common herpesvirus, CMV. Antibody levels were significantly different in PTSD *vs.* control subjects ($p = 0.016$). Finally, the authors used ANOVA to assess the contributions of comorbid diseases and found that only a small number of genes showed differential methylation across groups divided by comorbidity, indicating that the PTSD methylation findings are specific to PTSD and not affected by the increased rates of major depressive disorder or generalized anxiety disorder in PTSD populations.

5 Conclusions and future challenges

As stated in the Introduction, post-traumatic stress disorder is the only psychological disorder for which the onset of the pathological condition is fairly precise, *i.e.*, occurring after an extreme traumatic event. A wealth of often contradictory physiological findings point to the underlying complexity of the disorder. Although considered a psychological disorder, PTSD also affects the functioning of numerous non-neurological systems in the body, including the immune, endocrine and cardiovascular systems. In order to understand the relationship between the progression of PTSD and these other systemic responses, specific details about the molecular mechanism must be unraveled. In particular, details analogous to the increased negative feedback identified in the HPA axis¹³⁵ must be identified to understand the cardiovascular and immunological changes that frequently occur in PTSD. However, identifying the relevant components in these systems is further complicated by the high rates of comorbid diseases (biological noise) and experimental artifacts from high-throughput experiments (experimental noise). Addressing these

challenges will require careful consideration and likely novel adaptations to current machine learning strategies, including the use (or development) of data integration techniques that can accommodate data from multiple biological length scales.

The complexity of the disorder has posed a great challenge to the identification of biomarkers for diagnostic, prognostic and therapeutic applications. The time dependent profile, different for each subject, further complicates the diagnosis. A recent clinical study⁸⁹ has shown three different trajectories of symptom progression: rapid-remitting (fast decrease of symptoms in one to five months), slow-remitting (symptoms decrease after 15 months) and non-remitting (persistent symptoms). In order to understand the three clinical trajectories from a biological perspective and to explore the prognostic and therapeutic opportunities, it is increasingly evident that integration of both the biological and the clinical manifestations of PTSD is essential. Additionally, if these symptom trajectories or other distinct subtypes of PTSD exist, the choice of how and where these subgroups should be integrated or separated for biomarker identification analysis must be systematically explored. Specifically, the biological mechanism of PTSD development must be extensively studied (using longitudinal data) in all potential subtypes of PTSD to determine if overlapping signals exist. In the absence of overlapping biological mechanisms, multi-class classification strategies may be required. Since PTSD originates in the brain, it is extremely difficult to collect samples for scientific study. Alternatively, animal models are used to correlate with clinical findings, thus enabling one to draw inferences that may lead to diagnostic and therapeutic solutions.

Many animal models that exhibit one or more “PTSD-specific” symptoms have been proposed, including predator-exposure,^{145–147} exposure to single prolonged stress,¹⁴⁸ and exposure to foot shock^{149,150} with additional stress. However none to date have been widely accepted as an ideal model for PTSD. These models approximately mimic physiological and tissue-level responses in human PTSD subjects and have helped in identifying biological processes, such as the HPA axis, neurotransmitter receptor system, and others (see review by Pitman *et al.*¹²), that may be involved in PTSD. In the model studies section of this review, we presented a study by Cho *et al.*¹⁴³ based on the social defeat mouse model¹⁴⁰ that identifies key short-term physiological and histological changes that occur in PTSD-like mice. These changes, including cardiac histopathology differences as well as increased weight gain, show similarities to established comorbid PTSD phenotypes of obesity and cardiovascular disease. In addition to gaining insight into development and progression of PTSD, animal models may also provide an avenue for pharmacological studies. For example, the social defeat mouse model responds to chronic (but not acute) exposure to antidepressant medication, which agrees with human PTSD responses. In general, these proposed animal models of PTSD have shown agreement with previously identified pharmacological, organismal and tissue-level responses in humans. However, to the best of our knowledge, no study to date has compared long-term animal model

response with human PTSD subjects. These long-term responses, as well as detailed cellular and molecular changes across all time scales must be carefully addressed in future studies to more conclusively identify how well animal model findings can be translated to understand PTSD in humans.

In light of the complexity of the PTSD disease mechanism, a systems approach integrating both the clinical findings and animal validation studies is necessary to understand the underlying mechanisms driving PTSD and its variants. Such an approach establishes a multidimensional research direction combining patients, published animal models and mathematical analysis tools. The goals of such an approach is to (1) identify the differentially expressed genomic, proteomic and metabolic drivers of PTSD; (2) identify approaches to diagnose different clinically observed PTSD trajectories; (3) identify possible biological risk factors, prognostic indicators, and therapeutic strategies; (4) identify the underlying molecular principles to facilitate detailed studies of relevant molecular networks. In order to achieve these goals there is a need to integrate heterogeneous experimental data (*e.g.*, molecular and neuroimaging data) from different studies by employing new or existing mathematical tools to uncover useful information from large noisy datasets (on the order of a million data points per subject). In this paper we outline the preliminary studies that pave the way for a multidisciplinary integrated systems biology approach, and highlight some current challenges to identifying and understanding the molecular underpinnings of post-traumatic stress disorder.

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