A new bioinformatic insight into the associated proteins in psychiatric disorders

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Abstract

Background: Psychiatric diseases severely affect the quality of patients' lives and bring huge economic pressure to their families. Also, the great phenotypic variability among these patients makes it difficult to investigate the pathogenesis. Nowadays, bioinformatics is hopeful to be used as an effective tool for the diagnosis of psychiatric disorders, which can identify sensitive biomarkers and explore associated signaling pathways.

Methods: In this study, we performed an integrated bioinformatic analysis on 1945 mental-associated proteins including 91 secreted proteins and 593 membrane proteins, which were screened from the Universal Protein Resource (Uniport) database. Then the function and pathway enrichment analyses, ontological classification, and constructed PPI network were executed.

Results: Our present study revealed that the majority of mental proteins were closely related to metabolic processes and cellular processes. We also identified some significant molecular biomarkers in the progression of mental disorders, such as HRAS, ALS2, SLC6A1, SLC39A12, SIL1, IDUA, NEPH2 and XPO1. Furthermore, it was found that hub proteins, such as COMT, POMC, NPS and BDNF, might be the potential targets for mental disorders therapy. Finally, we demonstrated that psychiatric disorders may share the same signaling pathways with cancers, involving ESR1, BCL2 and MAPK3.

Conclusion: Our data are expected to contribute to explaining the possible mechanisms of psychiatric diseases and providing a useful reference for the diagnosis and therapy of them.

Keywords: Psychiatric diseases, Bioinformatics, Molecular biomarkers, Therapy, Signaling pathways

Background

Psychiatric disorders are generally regarded as neuropsychological and neurobehavioral lesions, and the impaired ability to understand new or complicated information. It is estimated that more than 450 million people worldwide suffer from various mental disorders, among which major depression will be the most debilitating one by 2030 (World Health Organization 2010). Also, the appearance of psychiatric disorders tends to take place in childhood with increasing prevalence and incidence. In addition, males with intellectual disability were even found to have an elevated risk of leukemia, brain, stomach, corpus uteri and colorectal cancers (Sullivan et al. 2004). However, the etiopathogenesis of many psychiatric diseases is still unclear.

Two thirds of psychiatric patients take physical examination at primary care facilities. They always complain about the lack of energy as well as general aches and pains, so that their doctors usually focus on the organic disorders but neglect the mental problems. Moreover, most kinds of psychiatric disorders often present similar symptoms, which frequently delays the early evaluation of neurodevelopment status and disease progression. To this day, the diagnosis of psychiatric diseases has mainly depended on the daily living experience of patients’ families and the subjective expression of patients. There seems not to be available biological tests objectively...
assessing the association between clinical symptoms and underlying molecular mechanisms.

Recently, with the development of bioinformatic technologies, we can get further insights into the pathogenesis of psychiatric disorders from the perspective of cells, circuits, and pathophysiological processes. And the recent findings of proteomic biomarker tests have made it more reliable and effective for the diagnosis and treatment of psychiatric disorders. For example, C-reactive protein (CRP) has been used as a biomarker to predict depression, and the increased CRP expression level was identified as an independent risk factor for de novo depression in women (Lopresti et al. 2014; Wium-Andersen et al. 2013). Besides, cytokines, neopterin, malondialdehyde, isoprostanes, and serum S100B have also been reported to be reliable and sensitive biomarkers of mood disorder. (Howgren et al. 2009; Rybka et al. 2013; Maes et al. 2012; Chung et al. 2013; Milaneschi et al. 2013; Schroeter et al. 2013). Because multiple biological process overlap in the progression of psychiatric diseases, greater successes may be made on mental disease treatment by targeting the early biomarkers and the crucial signaling pathways. These successes should also be useful for exploring underlying mechanisms of mental disorders.

In this study, we aimed to explore the potential proteomic biomarkers and pathological mechanisms in psychiatric disorders with the help of bioinformatic tools. First, we performed a comprehensive bioinformatic analysis on the associated proteins from UniProt database to determine their significant characteristics. Then the functional enrichment analyses and constructed PPI networks were carried out to identify key genes, proteins and signaling pathways. Psychiatric disorders investigated in this study were referred to autism spectrum disorder (ASD), schizophrenia (SCZ), bipolar disorders, major depression, anxiety, phobia, affective disorder, obsessive compulsive disorder, personality disorder and mental retardation psychiatry. Proteins with altered expression which influence the physiological and pathological progression of mental disorders may be involved in certain pathways,. And once these pathways are made clear, it will help us know better about the pathogenic mechanisms and develop the targeted therapeutic strategies. Therefore, bioinformatic classifications should be utilized as useful references for identification and characterization of biomarkers, which will be expected to give a new direction to psychiatry research.

**Results**

**Psychiatric-disease-associated proteins were screened and classified**

We screened reviewed proteins from the UniProt database and collected 2317 proteins associated with mental disorders. The retrieved proteins from various psychiatric disorders were classified as follows: mental retardation (585 proteins), intellectual disability (425 proteins), cognitive disorder (94 proteins), autism (437 proteins), schizophrenia (413 proteins), bipolar disorder (219 proteins), depression (63 proteins), anxiety (8 proteins), phobia (51 proteins), obsessive compulsive disorder (7 proteins) and personality disorder (2 proteins). Because there are 372 proteins counted repeatedly among them, 1945 proteins associated with human psychiatric diseases were obtained finally.

**The associated proteins and signaling pathways were identified by bioinformatic analysis**

After the DAVID functional annotation clustering, a total of 1075 GO terms were found to be associated with psychiatric diseases proteins. The implicative pathways of GO (top 20) were shown in Table 1. The result demonstrated that mental-disorder-associated proteins mainly enriched in 2 GO categories including biological process (BP) and cellular component (CC). In the BP group, the majority (26%) of the proteins were concerned with metabolic processes, followed by cellular processes (21%). While the proteins in the CC group were comprised of cell part (43%), organelle (21%) and membrane (17%) (Fig. 1). And the top 5 pathways significantly involved in GO terms contained transmission of nerve impulse, neuron projection, synaptic transmission, learning or memory and behavior.

By performing the KEGG analysis for enriched proteins, we obtained 30 significant KEGG pathways with P value <0.05, and we showed the top 20 in Table 2. The top 5 significant pathways were listed as follows: Neuroactive ligand-receptor interaction, Calcium signaling pathway, Long-term potentiation, Long-term depression, Prostate cancer.

Analysis of tissue specific proteins revealed that most of the proteins were expressed in brain (49%), epithelium (15.6%) and placenta (17%). In addition, there are 4 proteins members of the VCX family expressed exclusively in testis.

**Secretory proteins and cell surface proteins were identified**

After retrieval of the “LOCATE” database, membrane organization analysis indicated that 1261 proteins were soluble proteins including 91 secreted proteins, and that 593 proteins were membrane proteins containing 76 type I proeins, 168 type II proteins and 349 multiple transmembrane protein.

**Hub proteins were identified by PPI network**

PPI networks were established by Cytoscape software including 329 nodes and 1194 edges based on STRING
database. Then, the degree of all nodes was calculated and the proteins whose node degree > 20 were shown as follows: POMC (proopiomelanocortin, Degree = 41), NPS (neuropeptide S = 38), ESR1 (estrogen receptor 1, Degree = 35), BDNF (brain-derived neurotrophic factor = 32), BCL2 (B-cell CLL/lymphoma 2 = 31), PTEN (phosphatase and tensin homolog, Degree = 29), AR (androgen receptor = 29), MAPK3 (mitogen-activated protein kinase 3, Degree = 27), HDAC2 (histone deacetylase 2 = 27), AVP (arginine vasopressin = 24), CRH (corticotropin releasing hormone = 24), COMT (catechol-O-methyltransferase, Degree = 22), HTR2A (5-hydroxytryptamine receptor 2A = 21), ACTB (actin, beta = 21), DPYD (dihydropyrimidine dehydrogenase = 21), MT-ND1 (mitochondrially encoded NADH dehydrogenase 1 = 21), DRD2 (dopamine receptor D2 = 21), CCK (cholecystokinin = 20), HTR7 (5-hydroxytryptamine receptor 7 = 20). These hub proteins were displayed in Table 3 and the associated PPI networks were shown in Fig. 2.

### Discussion

The diagnosis of psychiatric disorders always lack biological gold standards because of the complicated nosology. The huge phenotypic variability in population also bring great challenges to explore credible biomarkers in the fields of drug abuse and neuropsychiatry researches. However, with the advancement of bioinformatic technology, a plenty variety of proteins have been explored to define different progression stages of mental diseases. We made full use of the information of psychiatric-disease-associated proteins from the public databases and performed comprehensive bioinformatic analyses to determine sensitive biomarkers and significant signaling pathways. Our work is expected to provide new insights into the diagnosis of psychiatric diseases and investigate the underlying mechanisms. In future, more bioinformatic data of mental diseases should be needed to evaluate risks and benefits of specific therapeutic approaches and to enhance drug research and development.

In our study, 1945 mental-associated proteins were screened from the public database. Then we performed an integrated bioinformatic analysis to determine the biological relationship between these proteins and psychiatric diseases. Ontological analysis demonstrated that most of the proteins, which could be classified into various functional groups, may participate in different biological activities of brain. Functional clustering analysis also implied that these mental-associated proteins were significantly related to five GO terms: transmission of nerve impulse, neuron projection, synaptic transmission, learning or memory and behavior. We found transmission of nerve impulse was the most important term enriched in psychiatric diseases, which played a fundamental role in dendritic morphology and electrophysiological properties. And the second most important term

| Category              | Term                                | Count | P value   |
|-----------------------|-------------------------------------|-------|-----------|
| Biological process    | GO:0019226: transmission of nerve impulse | 91    | 2.89E−32  |
| Cellular component    | GO:0043005: neuron projection        | 88    | 8.81E−32  |
| Biological process    | GO:0007268: synaptic transmission   | 81    | 3.90E−30  |
| Biological process    | GO:0007611: learning or memory      | 50    | 6.30E−30  |
| Biological process    | GO:0007610: behavior                | 102   | 2.91E−29  |
| Cellular component    | GO:0030424: axon                    | 52    | 5.71E−24  |
| Cellular component    | GO:0042995: cell projection         | 114   | 9.57E−23  |
| Biological process    | GO:0007267: cell–cell signaling     | 102   | 1.18E−20  |
| Biological process    | GO:0044057: regulation of system process | 68    | 7.66E−20  |
| Biological process    | GO:0007612: learning                | 30    | 1.37E−19  |
| Cellular component    | GO:0045202: synapse                 | 72    | 1.38E−19  |
| Biological process    | GO:0031644: regulation of neurological system process | 46    | 3.97E−19  |
| Biological process    | GO:0042596: fear response           | 18    | 4.61E−19  |
| Biological process    | GO:0051969: regulation of transmission of nerve impulse | 44    | 3.08E−18  |
| Biological process    | GO:0050877: neurological system process | 153   | 6.80E−18  |
| Biological process    | GO:0030900: forebrain development   | 44    | 1.22E−17  |
| Cellular component    | GO:0030425: dendrite                | 45    | 1.25E−17  |
| Biological process    | GO:0050804: regulation of synaptic transmission | 41    | 3.77E−17  |
| Biological process    | GO:0033555: multicellular organismal response to stress | 23    | 4.65E−17  |
| Biological process    | GO:0032990: cell part morphogenesis | 57    | 5.83E−17  |
neuron projection is associated with the transcription of ribosome-bound mRNA (Cook-Snyder et al. 2015). More than 80 proteins were identified in neuron projection term, and these proteins may influence metabolic process or cellular process in mental disorders, such as STIL (SCL/TAL1 interrupting locus) and XPO1 (exportin 1). Besides, ALS2 (alsin Rho guanine nucleotide exchange factor) and SLC6A1 (solute carrier family 6 member 1), which were the most prominent proteins related to the majority of top GO terms, could be valuable for the diagnosis and targeted therapy of psychiatry. Many studies reported that common motor neuron diseases have a close relationship with mutations in ALS2, resulting in juvenile lateral sclerosis and infantile-onset ascending spastic paralysis (Sheerin et al. 2014; Eker et al. 2014). The neurotransmitter r-aminobutyric acid (GABA) transporter 1, encoded by SLC6A1 gene, was verified as the most prominent biomarker in neural signal transduction pathways (Berg and Geschwind 2012). Nicola reported that even a tiny deletion of SLC6A1 gene could result in intellectual disability and multiple congenital anomalies (Dikow et al. 2014).

Through analysis of KEGG pathway enrichment, we finally got 30 significant pathways. Most of these pathways can be classified into two categories: neural signal transduction and cancer pathways. The neural signal transduction contains neuroactive ligand-receptor interaction pathway, calcium signaling pathway, long-term potentiation (LTP) and depression (LTD), gap junction, lysosome, Alanine, aspartate and glutamate metabolism. The most important neuroactive ligand-receptor interaction pathway has been applied into the assessment of antipsychotic treatment and the analysis of neuropsychiatric disorder model (Adkins et al. 2012; Kong et al. 2015), which is enriched by TSPO, THRA, THRB, GABBR3, GRIK1, GABRB2, GRIK2, TRPV1, GABRB1, HTR6, OXTR. HRAS (harvey rat sarcoma viral oncogene homolog) was a predominant protein participating in 18 significant pathways. It may play key roles in psychiatric disorders with pathological alteration. Schwartz mentioned that HRAS functioned in synaptic long-term potentiation (LTP) and memory formation (Schwartz et al. 2013). It was demonstrated that the regulation of long-term potentiation is important for adult neurogenesis, synaptic plasticity, and learning and memory (Madroñal et al. 2010; Cooke and Bear 2010; Park et al. 2015; Jing et al. 2015). Therefore, HRAS is an ideal marker for psychiatric disorders diagnosis. While cancer pathways contained prostate cancer, thyroid cancer, chronic myeloid leukemia, melanoma, glioma, ErbB signaling pathway, and mTOR signaling pathway. These pathways remind us a fire-new angle to explore useful information from a mass of known proteins which were ignored before. MAPK3 is known as extracellular signal-regulated kinases (ERKs) and almost appears in all pathways. This gene always regulates a variety of cellular processes such as cell cycle progression, differentiation and proliferation. The overexpression of MAPK3 may indicate the pathogenesis of fear, anxiety and related psychopathological conditions.

After the membrane organization analysis, 91 secreted proteins were predicted, which could be candidate proteins for diagnosis of heritable mental diseases. For example, SLC39A12 (solute carrier family 39, member 12), a putative zinc transporter gene, is highly expressed in adult brains. It can encode ZIP12 (zinc transporter 12) protein which is required for neurulation and neuronal development (Chowanadisai et al. 2013; Chowanadisai 2014). Any other protein including SIL1 (nucleotide exchange factor) precursor protein, IDUA (iduronidase), APP (amyloid beta precursor protein), NEPH2 (kin of IRRE-like protein), AOF2 (amine oxidoreductase), and

![Fig. 1 Broad functional classification of selected mental disorders-associated proteins. a Biological processes, b cellular component](image-url)
WDR81 (WD repeat domain 81) proteins are expected to be diagnostic markers or therapeutic targets. And we found that 593 membrane proteins were transporters, receptors, enzymes and adhesion molecules, which were involved in nerve cell interactions, signaling transduction and energy metabolism.

Hub nodes have more complicated interactions with mental diseases compared with other proteins, implying that they play a crucial part in mental disorders (Langfelder and Mischel 2013). Therefore, identification of the hub proteins may enhance the assessment of disorders progression, neurodevelopment status, and therapeutic approaches. The following genes: POMC, NPS, ESR1, BDNF, BCL2, PTEN, AR, MAPK3, HDAC2, AVP, CRH, COMT, HTR2A, ACTB, DPYD, MT-ND1, DRD2, CCK, HTR7 were discovered as the genes that encode hub proteins in the PPI network. The POMC gene had a highest degree of 41, encoding the adrenocorticotropin hormone (ACTH) peptide, which brought our attention to major depressive disorders and the regulation of hypothalamic–pituitary–adrenal (HPA) axis function. POMC gene could be a valuable marker for assessing the response to antidepressant treatment (Chang et al. 2015). Furthermore, the polymorphisms of COMT gene are associated with sporadic schizophrenia and mood disorder, respectively (Li et al. 2015; Pandolfo et al. 2015). COMT inhibitors had been used to ameliorate symptoms of parkinson’s disease (Müller 2015). Therefore, POMC and COMT are both potential pharmacological targets for treating psychiatric disorders.

The NPS and BDNF genes also have multidirectional physiological activities in brain. Recent studies showed that NPS was associated with the regulation of food intake, fear, addiction and anxiety (Oishi et al. 2014; Slatery et al. 2015; Wegener et al. 2012). Dine et al. showed NPS application to mouse brain slices activated neurotransmission and plasticity in hippocampal CA3 and CA1 synapses (Dine et al. 2013). And the strong anxiolytic effect of NPS may be concerned with the increased dopamine releasing in medial prefrontal cortex (Lukas

| Term           | Description                                         | Count | P value       |
|----------------|-----------------------------------------------------|-------|---------------|
| KEGG:04080     | Neuroactive ligand-receptor interaction             | 54    | 2.15E–10      |
| KEGG:04020     | Calcium signaling pathway                           | 34    | 6.88E–06      |
| KEGG:04720     | Long-term potentiation                              | 18    | 3.01E–05      |
| KEGG:04730     | Long-term depression                               | 18    | 3.70E–05      |
| KEGG:05215     | Prostate cancer                                     | 19    | 3.37E–04      |
| KEGG:04540     | Gap junction                                        | 18    | 9.74E–04      |
| KEGG:04150     | mTOR signaling pathway                             | 12    | 0.003294      |
| KEGG:05012     | Parkinson’s disease                                 | 21    | 0.004578      |
| KEGG:05216     | Thyroid cancer                                      | 8     | 0.008593      |
| KEGG:05220     | Chronic myeloid leukemia                            | 14    | 0.008781      |
| KEGG:00250     | Alanine, aspartate and glutamate metabolism         | 8     | 0.012514      |
| KEGG:04012     | ErbB signaling pathway                              | 15    | 0.012769      |
| KEGG:05218     | Melanoma                                            | 13    | 0.014087      |
| KEGG:05214     | Glioma                                              | 12    | 0.014554      |
| KEGG:05200     | Pathways in cancer                                  | 40    | 0.015047      |
| KEGG:00563     | Glycosylphosphatidylinositol(GPI)-anchor biosynthesis| 7     | 0.015127      |
| KEGG:05010     | Alzheimer’s disease                                 | 23    | 0.016819      |
| KEGG:04916     | Melanogenesis                                       | 16    | 0.017224      |
| KEGG:04142     | Lysosome                                            | 18    | 0.017275      |

| Gene symbol | Degree |
|-------------|--------|
| POMC        | 41     |
| NPS         | 38     |
| ESR1        | 35     |
| BDNF        | 32     |
| BCL2        | 31     |
| PTEN, AR    | 29     |
| HDAC2, MAPK3| 27     |
| AVP, CRH    | 24     |
| COMT        | 22     |
| HTR2A, ACTB, DPYD, MT-ND1, DRD2 | 21 |
| CCK, HTR7   | 20     |
The BDNF gene encodes a few proteins of nerve growth factor families, involving the process of neurogenesis and the realization of neuroprotective functions. It was also found that BDNF played a key role in memory and learning organization and motor function. The expression of BDNF protein always decreases in patients with degenerative and vascular dementias, anxiety, affective and behavioral disorders (Levada and Cherednichenko 2015). Jehn et al. indicated that the low-level expression of BDNF was associated with cognitive impairment and short-term memory, coupled with the increased IL-6, which would be implicated in pathophysiology of depression patients (Jehn et al. 2015). NPS and BDNF could be used as neurobiological markers in pathological process, and the increased concentration may provide a potential novel treatment option.

Increased references in PubMed also provided us other hub proteins involved in cancer pathways, such as ESR1, PTEN, BCL2, AR and MAPK3. The down-regulation of BDNF-Akt-Bcl2 anti-apoptotic signaling pathway could be responsible for pathogenesis of autism. (Sheikh et al. 2010) Borges also reported that ERK1/2 signaling pathway effectively regulate depression, anxiety and emotion (Borges et al. 2015). Therefore, these potential targets involved in cancers may share the same signaling pathway with mental disorders. To ascertain the accurate roles of these genes may clarify pathological mechanisms and provide valuable information for psychiatric therapy.

**Conclusions**

This study analyzed mental-associated proteins out of the reviewed database to explore the molecular mechanisms of psychiatric diseases by using bioinformatic methods.
Our findings are expected to contribute to determining mental biomarkers, such as secreted proteins SLC39A12, SIL1, IDUA, APP, NEPH2, AOF2, and WDR81. And hub proteins, such as POMC, COMT, NPS, BDNF, also have the potential to be applied as targets for the diagnosis and treatment of mental disorders.

Methods

Screen of psychiatric diseases associated proteins
The UniProt (Release 2011_10, http://www.uniprot.org) was used to screen psychiatric diseases associated proteins. All human proteins were extracted from the UniProt database involving all annotations, such as “mental retardation,” “intellectual disability,” “autism,” “Schizophrenia,” “bipolar disorder,” “major depression,” “anxiety,” “phobia,” “affective disorder,” “obsessive compulsive disorder,” and “personality disorder.” Then the associated proteins were further filtered manually from the downloaded data.

Enrichment bioinformatic analysis
Protein IDs were submitted to DAVID (http://david.abcc.ncifcrf.gov/), a platform for functional analysis, such as gene ontologies, protein domains, and pathways. Of these, the enrichment analysis of GO categories includes biological process (BP), molecular function (MF), and cellular component (CC). The P value was computed by right-sided hypergeometric tests, GO terms were considered significant with P value <0.01, and KEGG pathways with P value <0.05. Protein IDs extracted from UniProt also were uploaded to PANTHER (www.pantherdb.org) to identify molecular function and protein class term through gene expression tools.

Analysis of the membrane organization
The LOCATE (http://locate.imb.uq.edu.au/), a curated database, was used to describe membrane organization of proteins. Uniprot IDs of mental illness proteins were uploaded to the LOCATE server to select secretory and cell surface proteins, which are promising biomarkers.

PPI network construction and hub proteins identification
The STRING database, a Search Tool to pre-compute global resource and evaluate protein–protein interactions information (von Mering et al. 2003), was used to construct PPI networks. And we primarily analyzed screened proteins of human psychiatric diseases by the UniProt database. Then, we performed analysis for PPI networks using Geniscape of Cytoscape software (version: 3.2.1) (Shannon et al. 2003). From a previous document on biological networks, numbers of PPI networks subjected to the scale-free attribution (Lamb et al. 2006). Therefore, connectivity degree was calculated by statistics and important nodes in PPI networks were obtained, namely hub proteins. Finally, the node degree >20 was selected as the threshold to obtain hub proteins in our study.

Abbreviations
CRP: C-reactive protein; ASD: Autism spectrum disorder; SCZ: Schizophrenia; BP: Biological process; MF: Molecular function; CC: Cellular component; POMC: Proopiomelanocortin; NPS: Neuropeptide S; ESR1: Estrogen receptor 1; BDNF: Brain-derived neurotrophic factor; BCL2: B-cell leukemia/Lymphoma 2; PTEN: Phosphatase and tensin homolog; AR: Androgen receptor; MAPK9: Mitogen-activated protein kinase 9; HDAC2: Histone deacetylase 2; AQP9: Arginine vasopressin; CRH: Corticotropin releasing hormone; COMT: Catechol-O-methyltransferase; HTR2A: 5-Hydroxytryptamine receptor 2A; ACTB: Actin, beta; DYPD: Dihydropyrimidin dehydrogenase; MT1-MOD1: Mitochondrially encoded NADH dehydrogenase 1; DRD2: Dopamine receptor D2; CCK: Cholecystokinin; HTR7: 5-Hydroxytryptamine receptor 7; STIL: SCL/TAL1 interrupting locus; XPO1: Exportin 1; ALS2: Albin rhodanine nucleotide exchange factor; SLC6A1: Solute carrier family 6 member 1; SLC39A12: Solute carrier family 39, member 12; ZIP12: Zinc transporter 12; SIL1: Nucleotide exchange factor; IDUA: Iduronidase; APP: Amyloid beta precursor protein; NEPH2L: Kind of IRRE-like protein; AOF2: Aminoxiduredoxase; WDR81: WD repeat domain 81.

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Competing interests
All of the authors declare no competing interests and no financial relationship with any commercial entity that has an interest in the subject of this manuscript.
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