The involvement of secondary neuronal damage in the development of neuropsychiatric disorders following brain insults

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Neuropsychiatric disorders are one of the leading causes of disability worldwide and affect the health of billions of people. Previous publications have demonstrated that neuropsychiatric disorders can cause histomorphological damage in particular regions of the brain. By using a clinical symptom-comparing approach, 55 neuropsychiatric signs or symptoms related usually to 14 types of acute and chronic brain insults were identified and categorized in the present study. Forty percent of the 55 neuropsychiatric signs and symptoms have been found to be commonly shared by the 14 brain insults. A meta-analysis supports existence of the same neuropsychiatric signs or symptoms in all brain insults. The results suggest that neuronal damage might be occurring in the same or similar regions or structures of the brain. Neuronal cell death, neural loss, and axonal degeneration in some parts of the brain (the limbic system, basal ganglia system, brainstem, cerebellum, and cerebral cortex) might be the histomorphological basis that is responsible for the neuropsychiatric symptom clusters. These morphological alterations may be the result of secondary neuronal damage (a cascade of progressive neural injury and neuronal cell death that is triggered by the initial insult). Secondary neuronal damage causes neuronal cell death and neural injury not only in the initial injured site but also remote brain regions. It may be a major contributor to subsequent neuropsychiatric disorders following brain insults.

Keywords: neuropsychiatric disorders, secondary neuronal damage, brain insults, traumatic brain injury, neurodegenerative diseases, clinical manifestations, histomorphology, pathophysiology

Neuropsychiatric disorders, also called mental disorders or emotional disorders, are conditions that affect cognition, emotion, behavior, and substantially interfere with the ability to function (1). Clinical manifestations of neuropsychiatric disorders include memory and cognitive deficits, consciousness and sleep disturbances, mental and emotional symptoms, somatoform symptoms, and psychomotor performance deficits. Neuropsychiatric disorders are 4 of the 10 leading causes of disability worldwide (2). About 26.2% of the U.S. adult population and 27% of the European Union adult population aged 18 and older, meet the diagnostic criteria for neuropsychiatric disorders at some point in their lives (3, 4). Four hundred fifty million people worldwide were diagnosed with some form of neuropsychiatric disorder in 2001 (5).

The mechanism underlying neuropsychiatric disorders has not been fully elucidated and still remains controversial. Several different factors may be involved in the occurrence and development of neuropsychiatric disorders. Psychological (violent attacks, being abused or neglected, marital separation, divorce, the sudden death of a loved one, being unable to relate to others, etc.), biological (genetics, prenatal damage, infections, exposure to toxins, brain defects or injuries, chemical imbalances, and substance abuse), and social/environmental (such as poor social and work relationships, family argument, dispute at work, loss of job, retirement, unemployment or underemployment, serious financial problems, poverty, socioeconomic deprivation, and substance abuse) factors are usually considered to be the triggers that cause most neuropsychiatric disorders (6–11). Research thus far does not clearly demonstrate how these triggers can cause neuropsychiatric disorders in the brain, nor what the cellular and molecular changes associated with the mental diseases are. The histomorphological basis of a neuropsychiatric disorder has been generally ignored.

Every disease has its own histomorphological features. A disease will not occur without morphological alterations associated with the disease in cells, tissues, and organs. When the cause and pathomorphology of a disease are poorly understood, people usually mythologize the disease and attribute the symptoms to social and cultural problems such as a sedentary lifestyle, social isolation, depressed mood, economic crisis, overindulgence in sex, obesity, heredity, etc. For example, tuberculosis had been considered as a fatal wasting disease (consumption) caused by vampires before Mycobacterium tuberculosis was discovered in 1882 (12). Therefore, definition of morphological alterations at the cellular and molecular levels is a very important step in the diagnosis and
Table 1 | The initial causes of brain insults.

| Brain insult                    | Cause of injury                                                                                   |
|---------------------------------|--------------------------------------------------------------------------------------------------|
| Stroke                          | Progressive neuron death caused by disturbance in the blood supply to the brain (ischemia or hemorrhage) |
| Brain tumor                     | A tumor (an abnormal growth of cells) within the brain                                             |
| **TRAUMATIC BRAIN INJURY (TBI)**|                                                                                                |
| Blast-induced TBI               | Progressive neuron death caused by blast shock waves                                              |
| Closed head injury              | Progressive neuron death caused by external physical force(s)                                    |
| **NEURODEGENERATIVE DISEASES**  |                                                                                                |
| Parkinson's                     | Greatly decreased activity of dopamine-secreting cells caused by progressive neuron death in the basal ganglia |
| Alzheimer's                     | Progressive neuron death and synapses loss in the cerebral cortex, limbic system, and the brainstem |
| Amyotrophic lateral sclerosis    | Progressive death of motor neurons in the cortex and the brainstem, characterized by proteinaceous inclusions in neuron bodies and axons |
| Huntington’s (Chorea)           | An autosomal dominant mutation in Huntington gene on chromosome 4 causes cellular accumulation of protein clumps, inducing cellular toxicity, and progressive neuron death primarily in the basal ganglia and the cortex |
| **POISONING WITH HAZARDOUS CHEMICALS** |                                                                                               |
| Organophosphate pesticides/chemical nerve agents | Cholinergic neuronal deficits induced by irreversible AChE inhibition                                |
| Chemical neurotoxic substances  | Progressive neuron death caused by neurotoxicity-induced ionic imbalance in neurons               |
| Alcohol abuse                   | Progressive neuronal damage resulting from the combination of prolonged ingestion of alcohol, which may involve abnormal release of neurotransmitters and prolonged stimulation of reward circuitry |
| Drug abuse                      | The release and prolonged action of dopamine and 5-HT within the reward circuit of the brain activate the reward system and lead to drug addiction, which cause chronic neuropathy in the brain |
| **INFECTIONS BY PATHOGENIC MICROBES** |                                                                                               |
| HIV/AIDS                        | Progressive neuron death caused by the harmful inflammatory cascades including activation of macrophages and microglia, release of inflammatory mediators, and induction of uncontrolled inflammation |
| PANDAS                          | An autoimmune-mediated response to a streptococcal infection, which produces antineuronal antibodies to erroneously destroy normal neurons in the basal ganglia |

Treatment of the disorder in general. A recent study has suggested that neuropsychiatric disorders triggered by psychological and social/environmental factors may involve a volumetric blood surge created by a sudden rise in cardiovascular pressure after exposure to an emotionally and psychologically traumatic event. The volumetric blood surge will move quickly through blood vessels to the brain, dramatically increasing cerebral perfusion pressure and causing damage to both very small cerebral blood vessels in the brain and to the blood brain barrier (BBB) (13). This indicates that brain microstructural damage could be the histomorphological basis of neuropsychiatric disorders triggered by psychological and social/environmental factors.

In the present study, a clinical symptom-comparing approach was employed to identify the similarities and differences among the neuropsychiatric manifestations related to 14 different types of brain insult. To explore the potential mechanisms that cause neuropsychiatric disorders, the correlation between neuropsychiatric manifestations and pathomorphology of the brain was analyzed.

**BRAIN INSULTS SELECTED FOR STUDY**

Neuropsychiatric disorders frequently occur after various brain insults such as closed head injury (CHI) (caused by external physical forces), blast-induced traumatic brain injury (TBI) (caused by blast shock waves), stroke (including ischemic and hemorrhagic stroke), poisoning with hazardous chemicals [caused by organophosphate (OP) pesticides, chemical nerve agents, chemical neurotoxic substances, alcohol, drugs, etc.), infection by pathogenic microbes (such as HIV/AIDS and PANDAS), brain tumors (including gliomas, meningiomas, pituitary adenomas, and nerve sheath tumors), and degenerative diseases [such as Parkinson’s, Alzheimer’s, amyotrophic lateral sclerosis (ALS), and Huntington’s]. Although these brain insults can induce many of the same neuropsychiatric disorders, their initial causes of injury are totally different (Table 1).

**ACUTE BRAIN INSULTS**

**Closed head injury**

Closed head injury is a type of TBI caused primarily by motor vehicle accidents, sporting activities, accidental falls, assaults, etc., where the skull and dura mater remain intact. CHI can range from mild to severe. CHI causes initial damage only to specific brain parts, the impact site or opposite side of the skull. Post-injury pathophysiological changes are usually observed in areas bordering the injured sites, and not in the entire brain (14, 15).
**Blast-induced TBI**

Blast-induced TBI is a unique type of brain injury that is caused by blast shock waves in the victims who are exposed to a blast but do not sustain penetrating and blunt impact injuries. The vast majority (~90%) of blast-induced TBI cases are mild. The major mechanism of blast TBI may involve large-scale BBB damage and tiny cerebrovascular insults in the entire brain, which are caused by the blood surge moving quickly through large blood vessels to the brain from the torso (16).

**Stroke**

Stroke is a brain damage characterized by the sudden disruption of blood supply to an area of the brain and the rapid loss of brain function. It is the second leading cause of death worldwide. Stroke can be caused by an obstruction in the blood flow (ischemic stroke) or the rupture of a blood vessel that supplies the brain (hemorrhagic stroke). A loss of blood supply to part of the brain will initiate a pathophysiological cascade that leads to irreversible death of neuronal cells and permanent neurological dysfunction. The symptoms depend on the area of the brain affected, but generally symptoms include sudden loss of the ability to speak, memory and cognitive deficits, visual field defect, and hemiplegia (17, 18).

**Organophosphate pesticides/chemical nerve agents**

Organophosphate compounds are a group of highly toxic chemicals that have been used widely as pesticides and developed as chemical warfare nerve agents such as soman (GD), sarin (GB), tabun (GA), cyclosarin (GF) and VX. OP pesticides/chemical nerve agents disrupt the functioning of cholinergic nervous system by irreversibly inhibiting acetylcholinesterase (AChE). The inhibition of AChE by OPs results in both the accumulation of ACh at synapses of the central and peripheral nervous systems and overstimulation of cholinergic receptors that exceeds normal physiological limits. Acute, excessive stimulation of cholinergic receptors (mAChR, in the brain) causes cholinergic neuronal excitotoxicity and dysfunction, which are largely responsible for the cholinergic crisis in the acute phase of the OP exposures, and could subsequently cause brain damage and chronic neuropsychiatric consequences (19).

**Chemical neurotoxic substances**

Chemical poisoning is a condition in which a harmful chemical causes damage to human cells, tissues, or organs, usually by chemical reaction or other activity at the molecular level after a sufficient amount is absorbed by the human body. Chemical poisoning is a major public health concern and the second leading cause of injury and death worldwide (20, 21). Chemical poisoning can be caused by approximately 3,000 hazardous chemicals.

Chemical neurotoxic substances (such as acetone, benzene, lead, mercury, and strychnine) cause damage to the brain by disrupting ionic balance in neuronal cells, interrupting neural signal transmission between neurons, altering normal brain function, and inducing progressive neuronal cell death and neurodegeneration. Clinical manifestations may vary among the victims who are exposed to chemical neurotoxic substances, depending on the specific compound, the amount, the route, and the length of time of exposure, as well as on the age and health status of the person exposed (22, 23).

**HIV/AIDS**

The brain can be infected by pathogenic microbes such as bacteria and viruses. Infections by pathogenic microbes can cause harmful inflammatory cascades in the brain, because cerebral cellular infiltration in response to microbial invasion is weaker, and delayed and inflammatory response in the brain is much stronger and spreads more rapidly than in other organs or tissues (24). Therefore, an overactive immune response to microbial invasion in the brain should be the major contributor to infection-related brain damage.

HIV/AIDS is a disease of the human immune system caused by infection with HIV. Although the immune system is the principal target of HIV, the brain can be also affected by HIV. HIV-associated psychiatric disorders have been observed in more than 50% of patients with HIV/AIDS (25). These neuropsychiatric disorders may be caused by progressive neuronal cell death resulting from the harmful inflammatory cascades, including activation of macrophages and microglia to release inflammatory mediators such as cytokines, eicosanoids, and complements, and overstimulation of the immune system to lead to uncontrolled inflammation in the brain (26, 27).

**Pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections**

Pediatric autoimmune neuropsychiatric disorders associated with streptococcal infections (PANDAS) are neuropsychiatric disorders (usually obsessive–compulsive disorder (OCD) and/or tic disorders) following a Group A beta-hemolytic streptococcal infection (GABHS) in children (28, 29). The mechanism behind PANDAS may involve an autoimmune-mediated response to a streptococcal infection, which produces antineuronal antibodies to erroneously destroy normal neuronal cells in the basal ganglia (30).

**CHRONIC BRAIN INSULTS**

**Brain tumor**

Brain tumor is an intracranial solid neoplasm that is formed by an accumulation of abnormal growth cells. The most common types are glioma and astrocytic tumors from abnormal growth glial and astrocytes cells. The second most common type of brain tumor is meningial tumor that forms in the meninges. Pituitary adenomas are the third most common type of brain tumor; these are noncancerous tumors of the pituitary gland. Nerve sheath tumors are made up primarily of the myelin surrounding nerves and account for approximately 8% of all brain tumors. The symptoms of brain tumors mainly depend on tumor size and tumor location, but common clinical consequences include symptoms of intracranial hypertension, neurological and neuropsychiatric disorders, and psychomotor performance deficits (31–34).

**Alcohol abuse**

Alcohol abuse is an addictive illness that is caused by uncontrolled consumption of alcoholic beverages. Prolonged ingestion of alcohol will affect brain function by altering levels of neurotransmitters (GABA, glutamate, and dopamine) and inducing prolonged stimulation of reward circuitry, thus causing progressive brain damage. The brain areas that are commonly affected by alcohol are the cerebral cortex, cerebellum, hypothalamus and pituitary, and medulla.
In cerebral cortex, alcohol depresses the behavioral inhibitory centers, inhibits the thought processes, and slows down the processing of information from the eyes, ears, mouth, and other senses. This will cause the slowed reaction times, impaired memory, blurred vision, slurred speech, etc. If alcohol affects the cerebellum (the center of movement and balance), it leads to the “staggering gait of a drunk” or difficulty walking. Alcohol depresses the hypothalamus and pituitary to increase sexual urge, but decrease actual sexual performance. The hypothalamus and pituitary control sexual arousal and performance by coordinating automatic brain functions and hormone release. By acting on the medulla, alcohol induces sleepiness and deficits of consciousness, slow breathing, and decreases body temperature, which may cause death (35, 36).

Drug abuse
Drug abuse, also known as substance abuse, is a disorder that is caused by using an illicit drug to feel happy and excited, to be more productive or to help one stay awake, to improve athletic performance, or to ease another mental and emotional problem (such as stress, anxiety, or depression) (37, 38). Drug abuse has led to significant physical, psychological, and public health problems in almost every country. About 200 million people around the world use illegal drugs every year. More than 22 million Americans aged 12 and older (nearly 9% of the U.S. population) used illegal drugs in 2011 (39). The mechanism by which illicit drugs cause brain damage remains unknown. Some drugs may cause the neuronal cells to release abnormally large amounts of dopamine and 5-HT and produce prolonged enhancement of synaptic transmission within the reward circuit of the brain, thus causing progressive brain damage (40, 41).

MDMA (or called ecstasy) is an empathogenetic drug of the phenethylamine and amphetamine classes of drugs. The effects of MDMA on human brain are complex. MDMA may cause the neuronal cells to release abnormally large amounts of dopamine and 5-HT and produce prolonged enhancement of synaptic transmission within the reward circuit of the brain, thus causing progressive brain damage (40, 41).

Amyotrophic lateral sclerosis
Amyotrophic lateral sclerosis is a degenerative motor neuron disease characterized by the gradual degeneration and death of motor neurons in the cortex and the brainstem. ALS causes rapidly progressive weakness, muscle atrophy, fasciculations and spasticity, dysarthria, dysphagia, and dyspnea. Some patients with ALS have cognitive and memory deficits, depression, and other mental and emotional disorders (59, 60).
Huntington's disease
Huntington's disease (HD), also called Huntington's chorea, is an inherited neurodegenerative disorder that is caused by an autosomal dominant mutation in Huntington gene on chromosome 4. The Huntington gene mutation causes cellular accumulation of protein clumps, inducing cellular toxicity and progressive cell death primarily in the basal ganglia and the cortex (61–63). HD affects the individual's muscle coordination, judgment, memory, and other cognitive function, thus leading to uncontrolled movements, cognitive and memory deficits, and other neuropsychiatric disorders (64–66).

COMPARISON OF NEUROPSYCHIATRIC MANIFESTATIONS OF BRAIN INSULTS
Clinical manifestations of neuropsychiatric disorders include both signs and symptoms. The neuropsychiatric signs and symptoms indicate the presence of disease or abnormality in the brain. Some signs and symptoms (such as fatigue, nausea, vomiting, and hypersexuality) may be non-specific and can occur in many acute and chronic medical conditions, whereas most signs and symptoms are fairly specific for a brain disease or injury (such as memory loss, cognitive impairments, consciousness loss, anxiety, depression, and psychosis). The specific signs and symptoms should be meaningful and significant in assisting the diagnosis of brain insult(s).

The clinical symptom-comparing approach that was employed in the present study is a clinical research method to determine the same signs and symptoms that are commonly shared by two or more brain insults. Through literature search and review, the signs and symptoms relating to these brain insults are identified and categorized into different categories. All signs or symptoms identified are compared among different types of brain insults using Microsoft Excel. The same sign or symptom will be identified if this sign or symptom is commonly presented in all types of brain insults. A meta-analysis will be used to determine whether the same signs or symptoms exist in these brain insults and what the effect size of the same signs or symptoms is.

LITERATURE SEARCH ON NEUROPSYCHIATRIC SIGN/SYMPTOM CLUSTERS OF BRAIN INSULTS
Articles relating to the neuropsychiatric signs and symptoms of the 14 types of acute and chronic brain insults were searched between the years 1970 and 2013 using NIH PubMed database for keywords such as "Stroke," "Patient," and "Memory loss." A neuropsychiatric sign or symptom that was reported or described in at least three different articles was selected as a common sign or symptom for the brain insult. Fifty-five common signs or symptoms were identified for the 14 types of brain insult (Supplementary Table 1). They are categorized into five major categories: memory and cognitive deficits (memory loss, concentration difficulties, cognitive impairments, learning disabilities, etc.); consciousness and sleep disturbances (sleep disorder, insomnia, consciousness loss, etc.); mental and emotional symptoms (anxiety, depression, aggression, moodiness, etc.); somatoform symptoms (headache, reduced visuo-spatial abilities, blurred vision, fatigue, etc.); and impaired psychomotor and neuromotor functions (psychomotor retardation, lack of motor coordination, difficulty balancing, seizures/tremors, etc.) (Table 2).

IDENTIFICATION OF NEUROPSYCHIATRIC SIGNS/SYMPTOMS COMMONLY SHARED BY ALL OF THE 14 BRAIN INSULTS
Based on the 528 supplementary references in the Supplementary Material, 40% (22) of the 55 neuropsychiatric signs and symptoms were found to be commonly shared by all of the 14 brain insults, whether traumatic, infectious, toxic, oncogenic, or degenerative (Table 2). In the category "memory and cognitive deficits," six symptoms (memory loss, concentration difficulties, cognitive impairments, learning disabilities, forgetfulness, and impaired attention) are shared; in the category "consciousness and sleep disturbances," two symptoms (sleep disorder and insomnia) are shared; in the category "mental and emotional symptoms," six symptoms (anxiety, depression, aggression/agitation, psychosis, moodiness, and increased emotional sensitivity) are shared; in the category "somatoform symptoms," four symptoms (headache, reduced visuo-spatial abilities, blurred vision/visual field impairment, and fatigue) are shared; and in the category "impaired psychomotor and neuromotor functions," four symptoms (psychomotor retardation, lack of motor coordination, difficulty balancing, and seizures/tremors) are shared. Overlap of neuropsychiatric symptom clusters among different brain insults is shown in Figure 1. The 22 of the same neuropsychiatric manifestations were simultaneously identified in either acute (Figures 1A,B) or chronic (Figures 1C,D) brain insults. Many overlapping neuropsychiatric signs or symptoms were also found between each of the brain insults. This suggests that neuronal damage may be caused by these brain insults in the same or similar brain regions/structures or that the same or similar brain regions/structures can be possibly affected by different brain insults.

A meta-analysis was conducted to combine clinical manifestations of different brain insults and to determine the neuropsychiatric signs or symptoms commonly shared by all acute and chronic brain insults. The meta-analytic procedure adopted for this systematic study was based upon the methods described by Neyeloff et al. (67). The total numbers of patients presenting with all of the same neuropsychiatric signs or symptoms were used as the sample size for each brain insult. A matrix of sample-size weighted similarity coefficients between each of the 14 types of brain insults was computed in the meta-analysis. Forest plots of the meta-analysis give the effect estimate (the rate) and its 95% confidence interval for each insult. The overall combined effect size is indicated by a diamond. The meta-analysis of neuropsychiatric manifestations of the patients with brain insults shows that the same neuropsychiatric signs or symptoms exist in the 14 different types of brain insults (Figure 2).

IDENTIFICATION OF NEUROPSYCHIATRIC SIGNS/SYMPTOMS COMMONLY SHARED BY MORE THAN 10 DIFFERENT TYPES OF BRAIN INSULT
Other neuropsychiatric signs or symptoms were found to be commonly shared by more than 10 different types of brain insult: 6 neuropsychiatric signs or symptoms (impaired judgment, confusion, disorientation, fear/frightened, behavior stress,
Table 2 | The same neuropsychiatric signs and symptoms commonly shared by 14 types of brain insult.

| Brain insult                                      | Category of neuropsychiatric signs and symptoms                                                                 |
|--------------------------------------------------|---------------------------------------------------------------------------------------------------------------|
|                                                  | Memory and cognitive deficits | Consciousness and sleep disturbances | Mental and emotional symptoms | Somatoform symptoms | Impaired psychomotor and neuromotor functions |
| Acute brain insult                               |                                                                                |                                 |                            |                     |                                               |
| Closed head injury                               | Memory loss                     | Concentration difficulties         | Impaired attention          | Sleep disorder      | Lack of motor coordination                    |
| Blast-induced TBI                                | Concentration difficulties      | Cognitive impairments              | Learning disabilities       | Fatigue             | Difficulties in locomotion                     |
| Stroke                                           | Memory loss                     | Concentration difficulties         | Impaired attention          | Insomnia            | Difficulty walking and balance                 |
| OP/chemical nerve agents                         | Learning disabilities           | Cognitive impairments              | Impaired attention          | Anxiety             | Fatigue                                         |
| Chemical neurotoxic substances                   | Cognitive impairments           | Impaired attention                 | Learning disabilities       | Aggression/Agitation| Reduced visual-spatial abilities              |
| HIV/AIDS                                         | Impaired attention              | Learning disabilities              | Cognitive impairments       | Moodiness           | Lack of motor coordination                     |
| PANDAS                                           | Learning disabilities           | Cognitive impairments              | Impaired attention          | Psychosis           | Headache                                        |
| Brain tumor                                      | Impaired attention              | Learning disabilities              | Cognitive impairments       | Increased emotional sensitivity | Seizures, tremors, and lack of motor coordination |
| Alcohol abuse                                    | Impaired attention              | Learning disabilities              | Cognitive impairments       | Headache            | Lack of motor coordination                     |
| Drug abuse                                       | Impaired attention              | Learning disabilities              | Cognitive impairments       | Headache            | Lack of motor coordination                     |
| Parkinson’s                                      | Impaired attention              | Learning disabilities              | Cognitive impairments       | Headache            | Lack of motor coordination                     |
| Alzheimer’s                                      | Impaired attention              | Learning disabilities              | Cognitive impairments       | Headache            | Lack of motor coordination                     |
| Amyotrophic lateral sclerosis                    | Impaired attention              | Learning disabilities              | Cognitive impairments       | Headache            | Lack of motor coordination                     |
| Huntington’s (Chorea)                            | Impaired attention              | Learning disabilities              | Cognitive impairments       | Headache            | Lack of motor coordination                     |
| Chronic brain insult                             | Impaired attention              | Learning disabilities              | Cognitive impairments       | Headache            | Lack of motor coordination                     |
| The brain regions/structures may be involved in  |                                                                                |                                 |                            |                     |                                               |
| The limbic system (the hippocampus, the amygdala,|                                                                                |                                 |                            |                     |                                               |
| the cingulate gyrus, the thalamus, the hypothalamus,|                                                                                |                                 |                            |                     |                                               |
| the epithalamus, the mammillary body, basal ganglia system (the striatum), the cerebellum, and the cerebral cortex |                                                                                |                                 |                            |                     |                                               |
| The brain stem reticular formation, basal forebrain, hypothalamus, thalamus, and the cerebral cortex |                                                                                |                                 |                            |                     |                                               |
| The limbic system (the hippocampus, the amygdala,|                                                                                |                                 |                            |                     |                                               |
| the cingulate gyrus, the thalamus, the hypothalamus,|                                                                                |                                 |                            |                     |                                               |
| the epithalamus, the mammillary body, basal ganglia system (the striatum), the cerebellum, and the cerebral cortex |                                                                                |                                 |                            |                     |                                               |
| The cerebral cortex (dorsolateral prefrontal, insular, rostral anterior cingulate, premotor, parietal cortices, and parahippocampal gyrus) |                                                                                |                                 |                            |                     |                                               |
| The cerebral cortex (the posterior parietal, the primary motor, the premotor, and the supplementary motor cortices), the basal ganglia system, and the cerebellum |                                                                                |                                 |                            |                     |                                               |
FIGURE 1 | Overlap of neuropsychiatric symptom clusters among different types of brain insults. (A) Overlap of symptom clusters among closed head injury, blast-induced TBI, and stroke; (B) Overlap of symptom clusters among OP/chemical nerve agents, chemical neurotoxic substances, HIV/AIDS, and PANDAS; (C) Overlap of symptom clusters among brain tumor, alcohol abuse, and drug abuse; and (D) Overlap of symptom clusters among Parkinson’s disease, Alzheimer’s disease, ALS and Huntington’s disease.

and altered hearing and ringing in the ears) are shared by 13 brain insults; 2 neuropsychiatric signs or symptoms (dementia, and unusual temper tantrums/irritability) are shared by 12 brain insults; 4 neuropsychiatric signs or symptoms (mania, suicidal thoughts/Attempts, nausea, and vomiting) are shared by 11 brain insults; and 5 neuropsychiatric signs or symptoms (consciousness loss, restlessness, hallucinations, dizziness, and aphasia) are shared by 10 brain insults (Supplementary Table 1). Interestingly, the symptom of suicidal thoughts or attempts was commonly found in the cases of 11 different types of brain insult (CHI, blast-induced TBI, stroke, poisoning due to chemical neurotoxic substances, HIV/AIDS, alcohol abuse, drug abuse, Parkinson’s disease, Alzheimer’s disease, ALS, and HD). This suggests that suicidal thoughts or attempts may be a result of brain damage. A patient who has attempted suicide may be suffering from degeneration or death of neuronal cells in the brain, and need medical interventions for potential brain damage.

Post-traumatic stress disorder (PTSD) was found to be commonly shared by nine different brain insults (CHI, blast-induced TBI, stroke, poisoning due to OP pesticides/chemical nerve agents, poisoning due to chemical neurotoxic substances, HIV/AIDS, brain tumor, alcohol abuse, and drug abuse). PTSD is characterized by persistent re-experiencing symptoms, avoidance symptoms, and hyperarousal symptoms. It is closely associated with some brain insults such as TBI, stroke, brain tumor, and OP poisoning. PTSD symptoms developed in 50–80% of mild TBI patients (16), 25–50% of stroke survivors (68), 35% of childhood brain tumor survivors (69), and 8–23% of the victims of OP poisoning (70). PTSD is most likely to be a special delayed-onset neuropsychiatric disorder caused directly by brain injury or disease, not caused by exposure to traumatic stress.

CORRELATION BETWEEN CLINICAL MANIFESTATIONS AND HISTOMORPHOLOGICAL ALTERATIONS OF THE BRAIN
A neuropsychiatric disorder should have its morphological alterations in the brain. Any brain insult can certainly cause histomorphological, biochemical, or molecular biological alterations in some parts of the brain, such as the limbic system, basal ganglia system, the brainstem, the basal forebrain, the cerebellum, and the cerebral cortex. Unfortunately, these alterations at the cellular and molecular level are usually not easy to detect by using traditional diagnostic imaging techniques (e.g., ultrasound, X-rays, CT, and MRI), because tiny morphological changes (such as neuronal cell death or loss) in deep brain structures (such as the limbic system and basal ganglia system) can be often covered by other normal neural tissue layers of the brain. Much of the neuronal damage in
deep brain structures can be seen only under a microscope after death (13).

**MORPHOLOGICAL ALTERATIONS OF NEUROPSYCHIATRIC DISORDERS**

Recently, morphological alterations of some neuropsychiatric disorders (such as schizophrenia, depression, bipolar disorder, and drug abuse) have been verified in patients and animal models by using structural and functional brain imaging technologies and histopathological examinations. Reductions in brain volume have been reported in areas of the frontal cortex, temporal lobes, and brain stem in patients with schizophrenia (71–74). Morphological abnormalities in frontal–striatal–temporal areas (75), hippocampus (76), and the brain stem (72) have been observed in patients with depression and in a female monkey model of depression. Meta-analyses of structural MRI studies in bipolar disorder show that there is an increase in both the volume of the lateral ventricles and globus pallidus, and the rates of deep white matter hypointensities (77, 78). Functional MRI findings suggest that abnormal modulation between ventral prefrontal and limbic regions, especially the amygdala, likely contribute to poor emotional regulation and mood symptoms in the bipolar disorder patients (79). Morphological abnormalities in the brain stem have been observed in post-mortem tissue of bipolar disorder patients (72). In drug abusers, neuronal loss, neurodegenerative alterations, a reduction of glial fibrillary acidic protein-immunopositive astrocytes, and widespread axonal damage with concomitant microglial activation as well as reactive and degenerative changes of the cerebral microvasculature have been observed (80, 81). These evidences demonstrate that neuropsychiatric disorders can cause histomorphological damage in particular regions of the brain. Histomorphological damage in the brain will induce individual symptoms or clusters of neuropsychiatric disorders. The location, extent, and degree of brain tissue damage may correlate closely with the type and severity of neuropsychiatric symptoms.

**BRAIN REGIONS/STRUCTURES INVOLVED IN BOTH THE CATEGORY “MEMORY AND COGNITIVE DEFICITS” AND THE CATEGORY “MENTAL AND EMOTIONAL SYMPTOMS”**

Both “memory and cognitive deficits” and “mental and emotional symptoms” may involve neuronal damage in several parts of the brain including the limbic system (the hippocampus, the amygdala, the cingulate gyrus, the thalamus, the hypothalamus, the epiphysis, the mammillary body), basal ganglia system (the striatum), the cerebellum, and the cerebral cortex (frontal lobe and temporal lobe) (82, 83). The limbic system is a complex set of brain structures that lie on both sides of the thalamus, right under the cerebrum. It plays a key role in learning, cognition, long-term memory, emotion, motivation, social processing, and behavior. It also influences the endocrine system and the autonomic nervous system. Damage to the limbic system causes...
many neuropsychiatric disorders including anxiety disorder, bipolar (affective) disorder, psychopathic disorders, amnestic disorders, schizophrenia, and dementias (84, 85). The basal ganglia system is a group of nuclei located near the thalamus and hypothalamus. It consists of the striatum, the globus pallidus, the substantia nigra, the nucleus accumbens, and the subthalamic nucleus (86). It is strongly connected with the cerebral cortex, thalamus, and other brain areas to act as a cohesive functional unit by receiving input from the cerebral cortex and by sending outputs to the motor centers in the brain stem. The basal ganglia system is associated with a variety of functions, including reward learning involving dopamine release and transmission, cognitive and emotional functions, posture and movement, eye movements, and executive functions. Basal ganglia dysfunction has been implicated in a number of neuropsychiatric disorders such as Parkinson’s disease, HD, schizophrenia, Tourette syndrome, hemiballismus, and obsessive–compulsive disorder (87, 88). The cerebellum is a region that is located at the bottom of the brain underneath the cerebral hemispheres. It plays an important role in motor coordination, balance, posture, sensory perception, cognitive functions such as attention and language, and regulation of fear and pleasure responses (89). Cerebellar damage produces disorders related to fine movement and motor coordination (90). The cerebral cortex is the outermost layered structure covering the brain. It is often referred to as gray matter because it consists of cell bodies and capillaries. The cerebral cortex is the largest part of the human brain, associated with higher cortical functions such as cognition, memory, learning, attention, thought, language, information processing, abstraction, creativity, judgment, emotion, consciousness, voluntary muscle activity, somatic sensation, visual stimuli, and movement planning. Almost all neuropsychiatric disorders involve different degrees of damage that occur in the cerebral cortex (91, 92).

**BRAIN REGIONS/STRUCTURES INVOLVED IN THE CATEGORY “CONSCIOUSNESS AND SLEEP DISTURBANCES”**

Consciousness and sleep disturbances may be the result of neuronal damage in the brainstem reticular formation, basal forebrain, the limbic system (hypothalamus and thalamus), and the cerebral cortex (frontal lobe). The reticular formation is a region in the brainstem that has projections to the thalamus and cerebral cortex. It plays a central role in states of consciousness like alertness and sleep by regulating the sleep–wake cycle and filtering incoming stimuli to discriminate irrelevant background stimuli (93, 94). Injury to the brainstem reticular formation can result in coma, consciousness loss, and insomnia (95, 96). The basal forebrain is a group of structures located rostrally and ventrally to the striatum. It is considered to be the major cholinergic output of the brain, and can regulate wakefulness and REM sleep by the induction of ACh release. Damage to the basal forebrain can reduce the amount of acetylcholine in the brain and lead to impaired learning, amnesia, sleep disorder, and lethargy (97, 98). Other brain structures such as the hypothalamus and thalamus of the limbic system and frontal lobe of the cerebral cortex may also involve coordinated regulation of sleep–wake function and conscious states with the reticular formation and basal forebrain (99, 100).

**BRAIN REGIONS/STRUCTURES INVOLVED IN THE CATEGORY “SOMATOFORM DISORDERS”**

Somatoform disorders are a group of neuropsychiatric disorders characterized by physical symptoms that suggest physical illness or injury. However, the cause for the symptoms cannot be fully explained by a general physical illness because physical examinations and clinical lab tests do not indicate the presence of a physical illness (101). This has led to the hypothesis that the physical symptoms of these patients’ experience may be from a neuropsychiatric source associated with brain damage (102). Somatoform symptoms should not be the result of conscious malingering or factitious behavior. They may be relevant to the damage to some parts of the cerebral cortex (such as dorsolateral prefrontal, insular, rostral anterior cingulate, premotor, parietal cortices, and parahippocampal gyrus) (103, 104). Because the cerebral cortex controls sensory–motor coupling, sensory feedback system, and somatic sensation, cortical damage may cause increased sensitivity to internal physical sensations and pain. The increased sensitivity to somatic feelings may predispose patients to produce somatoform symptoms (105, 106).

**BRAIN REGIONS/STRUCTURES INVOLVED IN THE CATEGORY “IMPAIRED PSYCHOMOTOR AND NEUROMOTOR FUNCTIONS”**

Impaired psychomotor and neuromotor functions are defined as abnormalities of motor function that are associated with neuropsychiatric and neurological disorders. Psychomotor impairment (also known as psychomotor retardation) involves a visible slowing of movement (difficulty completing simple tasks), increased sensitivity to somatic feelings, and increased sensitivity to internal physical sensations and pain. The increased sensitivity to somatic feelings may predispose patients to produce somatoform symptoms (105, 106).

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**SECONDARY NEUROLOGICAL DAMAGE**

Unlike other tissues or organs, the brain is a very soft, tofu-like tissue. Although it is protected by the skull and cerebrospinal fluid and isolated from large or hydrophilic molecules (bacteria and
high-molecular-weight toxins) in the bloodstream by the BBB, the brain is extremely susceptible to injury and disease (115). An insult (injury or disease) to the brain may cause more serious consequences than to other tissues. Much of the damage done to the brain does not typically occur at the time of initial insult and does not result directly from the initial insult itself. The progressive deterioration in the brain functions following the initial brain insult results mainly from secondary neuronal damage, a cascade of progressive neural injury and neuronal cell death that is triggered by the initial insult. Secondary neuronal damage usually occurs in both the initial injured site and other brain regions remote from the initial injured site (such as the hippocampus, amygdala, thalamus, the brainstem, basal ganglia system, the cerebellum, and piriform cortex), and possibly continues in the hours, days, weeks, or months following the initial insult. It is largely responsible for the ultimate neuronal cell death and neural loss in the injured brain, and is a major contributor to subsequent neuropsychiatric impairments (19).

Secondary neuronal damage can be induced by the pathophysiological changes associated with the initial insults such as cerebral edema, inflammation, cytotoxicity, cerebral ischemia, reduced cerebral blood flow, cerebral hypoxia, hypercapnia, acidosis, altered neurotransmitter release, and raised intracranial pressure. Acute brain insults (such as TBI, stroke, poisoning with OP pesticides, chemical nerve agents or chemical neurotoxic substances, and infections by pathogenic microbes) can cause almost all pathophysiological changes as mentioned above. However, chronic brain insults (such as brain tumors, alcohol abuse, drug abuse, and neurodegenerative diseases) may induce only some of these pathophysiological changes. For example, a brain tumor leads to raised intracranial pressure, cerebral edema, inflammation, cerebral hypoxia, cytotoxicity, and altered neurotransmitter release. Neurodegenerative diseases, alcohol abuse, and drug abuse induce altered neurotransmitter release, cytotoxicity, cerebral edema, and inflammation. This may suggest that secondary neuronal damage can occur from the pathophysiological changes following various acute or chronic brain insults. The involvement of secondary neuronal damage will exacerbate neuronal injury and neurologic deficits, ultimately causing irreversible neuropsychiatric and neurological impairments (Figure 3).

**PATHOPHYSIOLOGICAL CHANGES ASSOCIATED WITH SECONDARY NEURONAL DAMAGE**

Cerebral edema, inflammation, cytotoxicity, loss of cerebral blood flow autoregulation, and the apoptosis program are thought to be the major contributors to secondary neuronal damage.

**Cerebral edema**

Cerebral edema is a condition characterized by an excess accumulation of water in the intracellular or extracellular spaces of the brain, causing raised intracranial pressure (116). It can be induced by all acute or chronic brain insults. Cerebral edema is closely associated with the development of secondary neuronal damage, and may be an early marker for progressive neural injury and neuronal cell death. The BBB disruption is the main cause of vasogenic cerebral edema, which is often caused by arterial hypertension, trauma, and tumor-induced release of vasoactive and endothelial substances. The BBB disruption allows intravascular proteins and water to enter and accumulate in the parenchymal extracellular space. Water in white matter can spread rapidly along fiber tracts to the gray matter and other remote regions of the brain, causing widespread cerebral swelling (117). Vasogenic cerebral edema is generally observed in acute brain insults (such as TBI, stroke, chemical poisoning, and infections by pathogenic microbes). Cytotoxic cerebral edema involves the intracellular accumulation of excess water, usually because of dysfunctioning sodium and potassium pumps in the neuronal cell membrane (118). It is often caused by chemical toxins, immune response to microbial infection, alcohol, illegal drugs, and damaged cells from trauma or neurodegenerative diseases. Cerebral edema causes many symptoms including headache, faintness, nausea, vomiting, and blurred vision. In some severe cases, it can lead to seizures and coma.

**Inflammation**

Inflammation is an immune response to harmful stimuli such as injury, microbial invasion, damaged cells, or foreign materials. Inflammation is a defense mechanism that protects the human body from infection and injury, and initiates the tissue-healing process. It leads to increased permeability of the vessels and an increased number of plasma proteins and leukocytes immigrating through vessel walls to the injured tissues (119). Activation of leukocytes results in the expression and release of classical inflammatory mediators including cytokines, eicosanoids, and complement, thus stimulating the signaling networks that regulate inflammatory responses to harmful stimuli (120). During the inflammatory process, destruction and healing of the tissue can happen simultaneously. If the inflammatory response is excessive in the injured tissue, tissue destruction induced by leukocytes will compromise the survival of the injured tissues or organs, causing more severe insults than the initial harmful stimuli. Inflammatory responses exist in almost all acute and chronic brain insults. In contrast to other tissues, neuroinflammatory response in the brain is induced and spread more rapidly because cellular infiltration in the brain in response to inflammation is weaker and delayed (121). When the brain is injured by the initial insult, it will activate microglial cells (the resident immune cells of the brain) to produce pro-inflammatory cytokines (such as interleukins, interferons, and growth factors), leading to a prolonged or exaggerated neuroinflammatory response in the injured areas of the brain (122). The exaggerated neuroinflammatory response is critical in the induction of progressive neural injury and neuronal cell death, which causes more severe tissue destruction and neuronal cell injury in the brain. Neuroinflammation has been recognized to be involved in the development of subsequent neuropsychiatric disorders following brain insults (123).

**Cytotoxicity induced by neurotransmitter imbalance**

Imbalances in some neurotransmitters (such as dopamine, 5-HT, ACh, and glutamate) in the brain can cause stimulation or inhibition of cell membrane receptors, leading to cytotoxicity. Cytotoxicity induces a variety of harmful effects in neurons, including altered calcium influx, free radical damage, oxidative stress, inflammatory responses, and apoptosis, potentially leading to neuronal...
Various acute and chronic brain insults cause post-injury pathophysiological changes including cerebral edema, inflammation, cytotoxicity, loss of cerebral blood flow autoregulation, and apoptosis in the brain. These pathophysiological changes induce secondary neuronal damage in the brain regions remote from the initial injured site (the limbic system, basal ganglia system, brainstem, basal forebrain, cerebellum, and cerebral cortex), leading to subsequent neuropsychiatric disorders.

Cytotoxicity involves the development of secondary neuronal damage triggered by many types of brain insult such as TBI, stroke, brain tumors, neurodegenerative diseases (Parkinson’s disease, Alzheimer’s disease, ALS, HD), and poisoning with hazardous chemicals (OP pesticides/chemical nerve agents, chemical neurotoxic substances, alcohol abuse, and drug abuse). Poisoning with hazardous chemicals induces over-release of some neurotransmitters (ACh, dopamine, 5-HT, and/or glutamate) from neurons to produce prolonged stimulation of post-synaptic neuron receptors, triggering excessive calcium release and increased calcium influx in neuronal cells (125). Increased calcium influx activates lipases, proteases, kinases, phosphatases, and endonucleases in potentially harmful metabolic cascades, thus arresting protein synthesis and depriving cells of enzymes or trophic factors essential to their survival (126). In addition, intracellular calcium overload can result in free radical-related damage and apoptosis by inducing excessive oxidative stress and inflammatory responses. Altered calcium influx can cause excitotoxic lesions in the affected neurons. Decreased release of neurotransmitters contributes to some neurodegenerative diseases such as Parkinson’s disease and Alzheimer’s disease. Greatly decreased activity of dopamine-secreting cells caused by progressive neuron death in the basal ganglia is the major cause of Parkinson’s disease (127). Reduced synthesis and activity of ACh in the cholinergic neurons may involve the onset of Alzheimer’s disease (128). Abnormal and decreased levels of glutamate transporters have been observed in ALS and Alzheimer’s disease. Inhibition of glutamate transporter causes elevated glutamate levels and alterations in glutamate receptors, ultimately resulting in excitotoxic lesion in neuronal cells (129).

Loss of cerebral blood flow autoregulation
Cerebral blood flow autoregulation is a process that maintains adequate and stable cerebral blood flow and blood pressure to the brain, and avoids underperfusion or overperfusion of the brain. It plays an important role in both delivering sufficient blood-containing oxygen and nutrients to the brain tissue for cerebral metabolic need, and removing CO₂ and other waste products from the brain. The autoregulation of cerebral blood flow is achieved primarily by dilation or contraction of cerebral arterioles under the influence of multiple complex physiological control systems (including carbon dioxide levels, cerebral metabolic rate, neural activation, activity of the sympathetic nervous system, posture, and other physiological variables) (130). Loss of the ability to autoregulate and control cerebral blood flow in cerebral arterioles occurs often in acute brain insults (especially in TBI and stroke), which results in massive cerebral edema, ischemia, and hypoxia. Impairment of cerebral blood flow autoregulation has been implicated in the development of secondary neuronal damage (131).
Apoptosis
In normal conditions, apoptosis generally confers benefits for the human body during the life cycle. Apoptosis can help eliminate old or dead cells by producing cell fragments (called apoptotic bodies). Apoptotic bodies are easily engulfed and quickly removed by phagocytic cells before they spill out onto surrounding cells and cause damage (132). Under the condition that a brain insult occurs, apoptosis becomes a programed cell death process that is fundamental to immune responses and is detrimental to tissue, which exacerbates brain damage and neurologic deficits (133). The apoptosis program can be triggered by excessive free radical formation, death-receptor (such as TNF and Fas receptors) ligation, DNA damage, loss of mitochondrial function, and lysosomal protease activation. Excessive apoptosis will result in uncontrolled cell proliferation, differentiation, and transformation, which plays a crucial role in triggering neuronal cell death (134, 135). Apoptosis, characterized by blebbing, cell shrinkage, nuclear fragmentation, chromatin condensation, and chromosomal DNA fragmentation, is a prominent feature in the brain after an insult (136). Apoptotic processes have been implicated in almost all acute and chronic brain insults, and may be the major contributor to ultimate neuronal cell death/loss in many brain regions remote from the initial injured site (e.g., the hippocampus, amygdale, thalamus, the brainstem, basal ganglia system, the cerebellum, and piriform cortex) (137, 138).

The pathophysiological changes associated with the initial brain insults trigger secondary neuronal damage in the brain. Secondary neuronal damage is the delayed and progressive neuronal cell death, neural loss, and axonal degeneration processes, which occur in not only the initial injured site but also other brain regions remote from the initial injured site. The mechanism by which secondary neuronal damage can happen in other brain regions remote from the initial injured site may involve multiple pathophysiological reactions that occur globally throughout the brain, including widespread cerebral edema and neuroinflammatory responses, loss of cerebral blood flow autoregulation, imbalances in neurotransmitters, defective neuronal signal transduction, altered ionic balance, and apoptosis. Secondary neuronal damage is the fundamental cause of subsequent neuropsychiatric and neurological disorders following brain insult. Neuropsychiatric disorders could be the long-term prominent clinical manifestations of the patients with brain insults, because chronic disability following brain damage is mainly expressed as mental sequelae rather than neurological deficits in humans (139). These chronic neuropsychiatric disorders are serious public health problems that result in the loss of many years of productive life and incur large healthcare costs.

CONCLUSION
Neuropsychiatric disorders are commonly induced by both acute (TBI, stroke, chemical poisoning, and infections by pathogenic microbes) and chronic (brain tumors, alcohol abuse, drug abuse, and neurodegenerative diseases) brain insults. Clinical manifestations of neuropsychiatric disorders may vary among the patients with different types of brain insult, depending on the cause, type, and severity of injury, acute or chronic condition, and the brain regions affected, as well as on the age and health status of the patient. However, 40% of neuropsychiatric signs and symptoms were found to be commonly shared by traumatic, infectious, toxic, oncogenic, and degenerative brain insults, suggesting that the same or similar brain regions/structures can be possibly affected by different acute and chronic brain insults.

The histomorphological basis for neuropsychiatric disorders might be the progressive neuronal cell death and neural injury in the brain regions that control memory, consciousness, sleep, and cognitive, emotional, somatic, psychomotor, and neuromotor functions, such as the limbic system, basal ganglia system, brainstem, basal forebrain, cerebellum, and cerebral cortex. Because most of the brain regions affected are far away from the initial injured site, the histomorphological alterations in the remote brain regions may mainly result from secondary neuronal damage triggered by the initial insult. Secondary neuronal damage may be largely responsible for subsequent neuropsychiatric disorders following various brain insults. However, it should be noted that clinical manifestations of neurological disorders may tend to be similar even when the histomorphological alterations in the brain region affected are dissimilar, because the brain has only a few common pathophysiological responses (such as cerebral edema, inflammation, cytotoxicity, loss of cerebral blood flow autoregulation, and the apoptotic) to injuries.

If a patient has neuropsychiatric signs and symptoms of unknown origin, physicians and psychiatrists should be aware that potential neuronal damage can possibly exist in the patient’s brain and not simply conclude that the patient has a psychological problem associated with social/environmental factors. Necessary treatment aimed at slowing down the process of neuronal damage and attenuating individual symptoms or clusters of neuropsychiatric and neurological disorders may help improve recovery of neurobehavioral functions. The clinical symptom-comparing approach used in the present study to identify the same clinical manifestations may be a useful tool for investigating the causes of neuropsychiatric disorders and other complex diseases of unknown origin.

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SUPPLEMENTARY MATERIAL
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