Towards a brain-based predictome of mental illness

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Abstract

Neuroimaging-based approaches have been extensively applied to study mental illness in recent years and have deepened our understanding of both cognitively healthy and disordered brain structure and function. Recent advancements in machine learning techniques have shown promising outcomes for individualized prediction and characterization of patients with psychiatric disorders. Studies have utilized features from a variety of neuroimaging modalities, including structural, functional, and diffusion magnetic resonance imaging data, as well as jointly estimated features from multiple modalities, to assess patients with heterogeneous mental disorders, such as schizophrenia and autism. We use the term "predictome" to describe the use of multivariate brain network features from one or more neuroimaging modalities to predict mental illness. In the predictome, multiple brain network-based features (either from the same modality or multiple modalities) are incorporated into a predictive model to jointly estimate features that are unique to a disorder and predict subjects accordingly. To date, more than 650 studies have been published on subject-level prediction focusing on psychiatric disorders. We have surveyed about 250 studies including schizophrenia, major depression, bipolar disorder, autism spectrum disorder, attention-deficit hyperactivity disorder, obsessive-compulsive disorder, social anxiety disorder, posttraumatic stress disorder, and substance dependence. In this review, we present a comprehensive review of recent neuroimaging-based predicors.

Abbreviations: AAL, automated anatomical labeling; ACC, anterior cingulate cortex; ADHD, attention-deficit hyperactive disorder; ANN, artificial neural network; AOD, auditory oddball; ASD, autism spectrum disorder; AUC, area under curve; AUD, alcohol use disorder; A2-CPT, AX version of continuous performance task; BOLD, blood-oxygen level dependent; BP, bipolar disorder; BPd, bipolar disorder depressed state; BR, bipolar disorder remitted state; cMDD, current MDD; CNN, convolutional neural network; Cohe-ReHo, coherence regional homogeneity; CSF, cerebrospinal fluid; CT, cortical thickness; DBM, deformation-based morphometry; DFM, discriminant function analysis; DLFPC, dorsolateral prefrontal cortex; DMN, default mode network; dMRI, diffusion MRI; DNN, deep neural network; DTI, diffusion tensor imaging; EC, effective connectivity; ED, Euclidean distance; EMD, earth mover's distance; Enet-TV, ElasticNet-total variation; FA, fractional anisotropy; FALFE, fractional amplitude of low frequency fluctuations; FC, functional connectivity; FFT, fast Fourier transform; FNC, functional network connectivity; GAD, generalized anxiety disorder; GBDT, Gradient Boosting Decision Tree; gLASSO, group least absolute shrinkage and selection operator; GLM, general linear modeling; GM, gray matter; GMD, gray matter density; GNB, Gaussian Naive Bayes; GPC, Gaussian process classifier; HC, healthy control; H-ELM, extreme learning machine; HOG-MHPC, fMRI histogram of oriented gradients-feature-based patient classification; ICA, independent component analysis; ITG, inferior temporal gyrus; KDA, kernel discriminant analysis; KNN, k-nearest neighbor; LDA, linear discriminant analysis; LDC, Linear discriminant classifier; LR, logistic regression; LRC, logistic regression classifier; MD, mean diffusivity; MD, mood disorder; MDD, major depressive disorder; MDDr, major depressive disorder remitted state; MDDo, major depressive disorder depressed state; MDR, major depressive disorder remitted state; MFMK-SVM, multi-kernel SVM learning method; ML, multi-layer perceptron; MNC, maximum norm clustering; mMLDA, modified maximum uncertainty linear discriminant analysis; MRS, magnetic resonance spectroscopy; MVPa, multivoxel pattern analysis; n.s., not specified; NMF, non-negative matrix factorization; NN, neural network; OCD, obsessive-compulsive disorder; PCA, principle component analysis; PLS, partial least squares regression; PTSD, posttraumatic stress disorder; RAVEN, regional analysis of brain volumes in normalized space; ReHo, regional homogeneity; RF, random forest; rMDD, remitted MDD; ROI, region of interest; sMRI, resting-state fMRI; RSN, resting-state network; RVM, relevance vector machine; SAM, social anxiety disorder; SAEs, sparse autoencoders; SD, substance dependence; SIFT, scale-invariant feature transform; SVM, support vector machine; SVM-RFE, support vector machine with recursive feature elimination; T2, T2*-relaxation; TRD, treatment-resistant depression; TSD, treatment-sensitive depression; VBM, voxel-based morphometry; VMHC, voxel-mirrored homotopic correlations; WBM, wavelet-based morphometry; WM, white matter; WMT, working memory task; WMV, white matter volume.

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1 | INTRODUCTION

With the first steps toward establishing modern psychiatry, and the first attempts to classify different mental phenomena and disorders into disease-specific categories, the need for diagnostic tools to allow for an objective evaluation of mental illness has surfaced. Current clinical diagnosis, disease evaluation and treatment plans for severe psychiatric disorders are solely based on cross-sectional self-reported clinical symptoms supported by information on longitudinal course and outcome. Researchers have been actively searching for objective, biologically based, disease indicators, or "biomarkers," and after several decades of experiments and attempts of classifying psychiatric disorders based on predefined symptom categories, we are currently at a turning point where a new paradigm has emerged: The Research Domain Criteria (RDoC; T. Insel et al., 2010). This approach aims at incorporating the most recent findings from clinical and genetic neuroscience, thereby opening the field to a dimensional approach informed by the specific neural pathophysiology underlying psychiatric disorders. By utilizing advanced neuroimaging techniques, it is now possible to study disease-specific structural and functional brain impairments. Neuroimaging modalities, such as magnetic resonance imaging (MRI), magnetoencephalography (MEG), and electroencephalography (EEG) offer tools to noninvasively study the neural structure of psychiatric disorders with exceptional accuracy. Using these powerful techniques, researchers have begun to understand the complex neural function and structure that may lead to specific disorders.

In recent years, there has been a growing trend in designing neuroimaging-based prognostic/diagnostic tools. As a result, there has been a lot of effort focused on the use of neuroimaging tools to automatically discriminate patients with brain disorders from healthy control (HC) or each other. Many of these studies have reported promising prediction performances with the claim that complex mental illness can be diagnosed robustly, accurately and rapidly in an automatic fashion. However, until now, these tools have not been integrated into the clinical realm. We believe a key reason for this is that many of the studies of this nature, despite the promising results on a specific research dataset, are not designed to generalize to other datasets, specifically the clinical ones.

In this systematic review, we surveyed the existing literature on the application of machine learning-based techniques for diagnostic-focused predictive analyses in psychiatric research and discuss current trends and future directions. While previous reviews have focused on a specific machine learning technique (Orru, Pettersson-Yeo, Marquand, Sartori, & Mechelli, 2012), a single disorder (e.g., schizophrenia [SZ], Calhoun & Arbabshirani, 2013; Kambeitz et al., 2015), major depression disorder (MDD; Gao, Calhoun, & Sui, 2018), and autism spectrum disorder (Retico, Tosetti, Muratori, & Calderoni, 2014)), a single imaging modality (B. Sundermann, Herr, Schwindt, & Pfeiderer, 2014), a small subset of disorders (Klüppel et al., 2012), or general brain-based disorders (Arbabshirani, Plis, Sui, & Calhoun, 2017), we aim to provide a comprehensive review of all major psychiatric disorders. So far, the most extensive review on major psychiatric disorders is the review article by Wolfers et al., where about 120 pattern recognition studies in SZ, mood disorders, attention-deficit hyperactivity disorder (ADHD), autism spectrum disorder (ASD), anxiety disorders, and specific phobias have been reviewed (Wolfers, Buitelaar, Beckmann, Franke, & Marquand, 2015). While there are some overlaps among the aforementioned studies and this current survey, to the best of our knowledge, this is by far the largest survey in the field of major psychiatric disorders based on the number of papers reviewed (about 250 papers). Further, in recent years, there has been an exponential growth of predictive analysis studies, and therefore, an updated survey is much warranted.

In this review, a general discussion of the current trends in the brain-based psychiatric "predictome" and their translational perspectives will be provided, along with highlighting some of the common challenges and guidelines for future directions. We also discuss emerging trends in neuroimaging such as data sharing, multimodal brain imaging, and differential diagnosis. The main goals of this study include: (a) to review and systematically compare a large number of recent MRI-based mental disorder diagnostic/prognostic studies in SZ, MDD, bipolar (BP) disorder, ASD, ADHD, obsessive–compulsive disorder (OCD), social anxiety disorder (SAD), posttraumatic stress disorder (PTSD) and substance dependence (SD), and (b) to discuss pitfalls and promises of existing machine learning techniques, and (c) to provide our vision and future directions to address some of the challenges. While there are a number of challenges remain to be addressed, brain-based predictome studies have made a considerable progress in recent years. We hope that, with more sophisticated machine learning approaches integrated with large-scale data, predictive modeling tools will transition from the "proof-of-concept" stage to the "ready for clinical implementation" stage in the near future.

2 | DEVELOPING A MENTAL ILLNESS PREDICTOME PIPELINE

Predictome studies using neuroimaging data aim to extract multivariate brain network features from one or more neuroimaging modalities to predict outcome measures such as specific psychiatric diagnoses.
Typically, after feature extraction and selection, a classifier is trained in a supervised or semi-supervised way with a predefined set of labels. Further model validation is performed either using an independent testing dataset or by incorporating a cross-validation (CV) scheme. Figure 1 presents the most common components of a brain-based predictome pipeline of mental illness prediction using neuroimaging data. While specific pipelines might vary at different preprocessing and postprocessing stages, conventional predictome analyses typically include the following steps: (a) feature extraction and selection/reduction, (b) classifier training, (c) classification and CV, and (d) performance evaluation.

### 2.1 Feature extraction, selection/reduction

The first step of a predictome analysis is to transform neuroimaging data into features (i.e., deciding what to use as features and extracting these feature values from the data). A neuroimaging feature refers to...
any derived variable containing valuable information about the class labels that can be extracted from the data.

In this survey, we reviewed and highlighted predictome studies based on the type of features used for classification purposes, including voxel-based, region-based, and brain-network based feature selection approaches. For example, features can simply be a set of brain voxels within a particular brain network, or a region of interest (ROI), multivariate data-driven (e.g., using independent component analysis [ICA]) brain networks, or jointly estimated multimodal features, as seen in Figure 1b. A voxel-based approach employs feature extraction at the brain voxel level, while a region-based approach identifies and extracts predefined region-of-interests (ROIs) as features based on a brain atlas (either functional or structural). A network-based feature extraction approach, such as ICA, aims at combining multiple voxels across brain network (Calhoun, Adali, Pearlson, & Pekar, 2001b; McKeown et al., 1998).

In addition to feature extraction, it is often important to reduce the number of features from high-dimensional neuroimaging data before proceeding with model training. In the context of neuroimaging, feature selection can help achieve higher accuracy rates (Ad-Dab’bagh et al., 2006), and allow a more specific focus on the underlying brain regions that account for between-group differences (Plitt, Barnes, & Martin, 2015). Indeed, the number of features in neuroimaging data is large with many irrelevant features not contributing to the prediction power of the model, and not all disorders affect every brain network in the same way. Thus, some brain-based features might not contribute to the diagnosis labels, and some features may capture redundant information already uncovered by other features. Computational time and model generalization can also be improved by excluding redundant and unrelated features (Dash & Liu, 1997; Guyon & Elisseeff, 2003; Moradi et al., 2015). Feature selection approaches (e.g., principle component analysis [PCA]) project the high-dimensional neuroimaging data into a lower dimensional space with a goal of preserving model discriminative power. Although not an essential step, in order to improve the strength of the prediction algorithm, it is important to select both optimum and meaningful features (Chu et al., 2012; Cuingnet et al., 2011). In a supervised learning approach, most discriminative features are selected to amplify the signal and reduce the noise. Often, prior information is used to address the dimensionality issue of neuroimaging data. Based on the characteristics of features and the type of learning problem, a particular feature selection approach is used (Mwangi, Tian, & Soares, 2014). Common feature selection approaches include: (a) expert feature selection (based on prior knowledge) and (b) automatic feature selection (based on a feature selection algorithm). A combination of these two approaches can also be used for feature selection. For example, an expert feature selection approach can first be implemented by selecting a previously known disorder-specific ROI, and then an automatic feature selection algorithm can be used to favor discriminative features within the predefined ROI. Note that, to avoid performance bias, feature selection and extraction methods should be limited to training dataset.

2.2 | Classifier training

A classifier is a function that takes features as input and generates a class label prediction. Based on the learning function and underlying assumptions, different types of classifiers can be developed. Neuroimaging studies have applied various classifiers for mental illness prediction. The dimensionality issue associated with the relatively large number of features and the small number of samples should be accounted for while applying such classification algorithms. Typically, the classifier learns a rule and separates the underlying classes optimally. Any type of classification or regression algorithm can be used for training purpose, such as linear and logistic regression algorithms, multilayer neural networks and Gaussian approaches (Bishop, 2006), in the current review, we have limited our focus on classifiers using discrete outcome measures (i.e., diagnostic labels), with the exception of discussion on translational perspective and advanced predictive modeling (Sections 5 and 6).

2.2.1 | Nearest-neighbor

The simplest form of classifier is known as the “nearest-neighbor” which does not require any explicit learning of a classification function. Using the nearest-neighbor approach, classification of an independent test sample is performed by identifying the most similar measures, for example, lowest Euclidean distance, between the training and testing samples, and then assigning the label of the training sample (i.e., nearest neighbor) to the test sample.

2.2.2 | Discriminative and generative models

Other classifiers that require explicit learning function can be categorized as discriminative and generative models (Trevor, Robert, & Friedman, 2009). A discriminative classifier directly learns to predict from the training data using a learning function based on predefined parameters. In contrast, generative classifier learns a statistical model to generate class labels by modeling the distributions of feature values that are conditional on example class labels.

2.2.3 | Support vector machine

During the training stage of a supervised learning, data labels are used to optimize the model by finding a hyperplane or decision boundary that can maximally discriminate between groups. The most common choice for a simple learning function is predicting class labels based on a linear combination of the features that might influence the outcomes. A linear classifier can be viewed as learning a line or boundary (i.e., decision boundary) that separates points within the two classes and discriminates their labels. For instance, a linear support vector machine (SVM) is such a classifier that learns the decision boundary. Due to its widespread use and promising results in neuroimaging-based
prediction, SVM is the most commonly seen classifier in our current survey. The SVM algorithm is typically intended for binary classification that aims at maximizing the boundary between different classes in a higher dimensional space. Mathematically, the discriminant function for SVM consists of a weight vector orthogonal to the decision boundary, and is specified by the data points that lie closest to the decision boundary, known as support vectors. This decision boundary further defines the classification rule of new, unseen cases.

### 2.2.4 Linear discriminant classifier

Another powerful linear model is the linear discriminant classifier (LDC) that attempts to separate classes by maximizing the variance of between-class to within-class ratio. An example of probabilistic discriminant model is the logistic regression classifier (LRC) that focuses on learning an optimum decision rule by modeling the log-odds ratio as a linear combination of predictor variables (i.e., feature). Both LDC and LRC methods yield probabilistic predictions that a new case can be assigned to a particular class and a class label.

### 2.2.5 Gaussian process classifier

Further, a Gaussian process classifier (GPC) is a probabilistic model and is a Bayesian extension of LRC (Wolfers et al., 2015). Briefly, GPC is first trained using the training feature to determine an optimized predictive distribution distinguishing between case and control. Note that, parameters relevant to this predictive distribution are estimated by maximizing the logarithm of the marginal likelihood on the training features. During the training stage, GPC then predicts the case and control by providing the predictive distribution of the test data using a sigmoid function (Frangou, Dima, & Jogia, 2017). For technical details of GPC, refer to Schrouff et al. (2013).

### 2.2.6 Neural network classifier

Also, artificial neural network classifiers (NNC) have recently become popular for modeling biological networks. Multilayer NNC is the extension of linear perceptron classifier, which can yield complex nonlinear decision boundaries. Typically, the structure of NNC includes an input layer, hidden layer(s) and an output layer. Neurons from each of these layers are connected to the neurons of the subsequent layers. A variety of nonlinear transfer functions of the hidden later neurons can be used (e.g., sigmoid function). Briefly, during the training phase, the weights across a set of artificial connected neurons are adjusted for learning purposes using backpropagation technique (Werbos, 1990), and then used for classification. For example, in case of mental illness prediction, artificial NNC analyzes the training labels (i.e., healthy versus disorder) and learns to identify a test example.

### 2.2.7 Random forest

Other recent and more powerful approaches for brain-based prediction include random forest and deep learning classifiers. In the random forest classifier, which is an ensemble of decision tree classifiers, multiple levels of randomization are integrated (Breiman, 2001). Using a randomized subset of the training data, each decision tree is grown, and each node is subsequently formed by searching through a random subset of training features. For each feature, the classifier estimates a score to highlight the feature’s discriminative power (i.e., Gini Importance [GI] score). The random forest approach offers improved generalization accuracy as it randomizes training subjects, particularly in cases with a relatively smaller training subjects compared to the number of training features. Further, the random forest classifier provides nonlinear decision boundaries, which helps to model nonlinear patterns of features during training.

### 2.2.8 Deep learning

Deep learning classifiers have recently become an attractive choice for mental illness prediction (Calhoun & Sui, 2016; Han, Huang, Zhang, Zhao, & Chen, 2017; lidaka, 2015; Jang, Plis, Calhoun, & Lee, 2017; J. Kim, Calhoun, Shim, & Lee, 2016; Plis et al., 2014). Deep learning classifiers can learn the features with optimal discriminating power directly from the raw data by using a hierarchical approach (Schmidhuber, 2015; Vieira, Pinaya, & Mechelli, 2017). This provides a great advantage over conventional classifiers that require explicit feature reduction steps. By applying nonlinear transformations to the raw data, deep learning classifiers automatically overcome issues with feature selection which is particularly helpful for higher-dimensional features or data with lack of prior knowledge.

### 2.3 Classification and CV framework

Once the classifier learns the decision rule based on features from a training set, the next step is to validate the model in a testing set. In order to mitigate performance bias and overfitting from predictive modeling, it is critical to analyze the training and testing datasets independently. During the training stage, the classifier learns to predict the labels from the training features based on the associated learning algorithm. For example, for learning problems without complex, iterative feature selection, the trained classifier is tested on previously unseen testing data (Wolfers et al., 2015). In order to achieve better model performance, a classifier should be trained with as much training data as possible, which is often a challenging issue in neuroimaging-based prediction studies. CV approach offers us to train classifiers with a higher number of training samples. A common CV approach is to repeatedly evaluate model performance using multiple training and testing partitions, a validation approach known as k-fold CV (k: number of data partitions; Kohavi, 1995; Patel, Khalaf, & Aizenstein, 2016). Other popular CV approaches include, leave-one-out
(LOO-CV), and holdout. LOO-CV is an iterative process, typically used on smaller sample size, where $k$ is equal to the number of samples and every subject in the whole sample is left out once for testing the classifier. Briefly, the LOO-CV procedure includes the following steps: (a) leave one sample out, train on the remaining ones, make a prediction for this sample (b) repeat for each sample in turn, and (c) compute the accuracy of the predictions made for all the samples. While a popular choice, leaving each sample out could become computationally expensive as it requires training of as many classifiers as the number of samples. In addition, LOO-CV has also been shown to potentially introduce some prediction bias (Varoquaux et al., 2017), as it could introduce high variance by providing more data during the training stage, which could also result in overfitting (Elisseeff & Pontil, 2003; Refaelzadeh, Tang, & Liu, 2009). Because of this, the preferred approach is $k$-fold CV where $k$ < number of samples. Common choices for partitioning are $k = 10$ or $k = 5$, corresponding to leaving out 10 or 20% of the total samples during each validation fold. Other important considerations for designing a CV procedure include: (a) inclusion of examples from all classes in the training data for better prediction accuracy, (b) having roughly equal number of samples across classes (i.e., balanced classes), and (c) inclusion of correlated samples in the same fold to avoid misleading performance that may accurately predict test samples with a correlated counterpart in the training set (Pereira, Mitchell, & Botvinick, 2009).

Performance measures, including accuracy, are averaged across iterations for the training and testing phase. For a supervised approach, a model is optimized using labeled data to find a discriminative decision boundary or hyperplane differentiating between case and control groups. The model parameters are optimized for maximum discrimination between groups. The CV approach helps ensure generalization of the training. During the classification stage, the trained model is then used to predict the label for new, unseen observations from testing set. For an unbiased generalization, it is important that the testing data do not overlap with the training data (Lemm, Blankertz, Dickhaus, & Müller, 2011). Further, the new data should be preprocessed in the same way as the training data.

More recently, another type of CV has been introduced, where various types of classifiers are cross-validated by running multiple classifiers on the same training data. For example, “Polysifier” can be used to cross-validate multiple classifiers, where the baseline is first computed by applying multiple classifiers, such as nearest neighbors, linear SVM, radial basis function or RBF-SVM, decision tree, random forest, logistic regression, naïve Bayes, and linear discriminative analysis (LDA; http://mialab.mrn.org/software/#polysifier).

### 2.4 Measures for performance evaluation

The most commonly used performance evaluation measures for predictive algorithms include accuracy, sensitivity, specificity, and the receiver operating characteristic (ROC) curve. These measures provide an evaluation of how accurately a classifier can generalize to new test samples (i.e., cases). In a clinical context, accuracy indicates how accurately the model classifies the cases and controls, sensitivity shows the proportion of true positives correctly identified (i.e., what percentage of cases are truly identified), and specificity demonstrates the proportion of true negatives correctly identified (i.e., what percentage of controls are truly identified) by the model. The overall performance of the model can be assessed by the ROC curve which provides a summary of the area under the curve (AUC). A high sensitivity suggests that only a few participants are falsely diagnosed as HCs while actually being patients, and a high specificity indicates that a few participants are falsely diagnosed as patients while actually being HCs. The accuracy refers to the total proportion of samples correctly classified. The ROC curves show the balance between the true positive rate (sensitivity) and the false positive rate (1-specificity) across a range of decision thresholds within the model. To avoid bias by potential imbalances between groups, a common practice is to report balanced accuracy measures, by taking an average accuracy obtained for each diagnostic label (Brodersen, Ong, Stephan, & Buhmann, 2010).

A useful measure to summarize the classification performance is to provide a confusion matrix, which represents actual labels on one side and the predicted labels on the other side. This is more important with models predicting more than two groups (Baldi, Brunak, Chauvin, Andersen, & Nielsen, 2000). Other useful performance measures can be extracted from the confusion matrix including positive predictive value (PPV), negative predictive value (NPV), F1-score (harmonic mean of precision and recall), and G-mean (geometric mean of precision and recall; Alberg, Park, Hager, Brock, & Diener-West, 2004). Positive and negative predictive values are important for predictive studies as they directly quantify the potential utility of the classifier for clinical diagnosis. The positive predictive value is defined as the number times the classifier correctly predicted participants as patients (i.e., positive diagnosis) divided by the total number of positive predictions. The negative predictive value is defined as the number of times the classifier correctly predicted a negative diagnosis divided by the total number of negative predictions.

### 3 Prediction of mental illness using neuroimaging techniques

With recent advancements in medical imaging technology, neuroimaging data is being collected more rapidly and at finer resolution than ever before. In recent years, there has been an increasing interest in leveraging this vast amount of brain data across analytic levels, acquisition approaches, and experimental designs to achieve a deeper understanding of brain structure and function. In this review, we use the term “predictome” to describe the use of multivariate brain network features from one or more neuroimaging modalities to predict mental illness. In the predictome, multiple brain network-based features (either from the same modality or multiple modalities) are incorporated into a predictive model to jointly estimate features that are unique to a disorder and predict subjects accordingly. Here, we review recent predictomic approaches used for neuroimaging classification and prediction, and provide an overview of studies for prediction of mental illness from their healthy counterparts.
3.1 | Survey procedure for the current literature review

The current review is based on a comprehensive literature search for research articles performing MRI-based predictive analyses of psychiatric illnesses. A systematic literature search was performed primarily in PubMed from 1990 to 2018, and more than 550 articles were found. SZ (Calhoun, Kiehl, Liddle, & Pearlson, 2004) was one of the first disorders investigated with predictive analyses, followed by major depressive disorder (MDD; Fu et al., 2008; Marquand, Mourão-Miranda, Brammer, Cleare, & Fu, 2008) and BP disorder (Arribas, Calhoun, & Adali, 2010). ADHD (C.-Z. Zhu et al., 2008), ASD (Ecker et al., 2010), PTSD (Q Gong et al., 2014), OCD (Weygandt et al., 2012), SAD (Liu et al., 2015) and SD (Vergara, Mayer, Damaraju, Hutchison, & Calhoun, 2017; Vergara, Weiland, Hutchison, & Calhoun, 2018). Figure 2 illustrates the systematic literature search process for this current study. Briefly, the search consistent of the following steps: (a) different terms related to classification/machine learning as well as their abbreviations (e.g., support vector machine, search with the term “SVM”), (b) all terms and abbreviations related to structural, functional and diffusion MRI (dMRI) combined with the term “biomarker”, and (c) all terms and abbreviations for one of the eight psychiatric disorders mentioned above. These steps were repeated for all disorders, and the identified references were further checked for missed publications which were included in the review as well. An additional screening process included on the relevance of the publications for the current review. Finally, we focused on all publications using a predictive analyses approach on MRI-based data in a case–control design of mental illness diagnoses that explicitly evaluated classification performance measures (e.g., overall classification accuracy). Further, the same search procedure was repeated in Google Scholar in order to reduce the probability of missing relevant articles of interest. About 250 papers were eventually selected for this survey that includes: 101 SZ, 61 MDD/BP, 35 ADHD, 38 ASD, 1 PTSD, 12 OCD, 2 SAD, and 7 SD. We categorized these articles based on a scheme developed for this review as depicted in Figure 1a–e and a summary of all articles is presented in Tables 1–8. Further, we limited our search range to journal articles in English published up until December 2018. Search criteria also included exclusion of articles without available full-text, and similar papers published by the same authors. For each study, key aspects such as imaging modality, classification method, sample size, and type features were investigated in a quantitative manner, as seen in Figures 3–6.

3.2 | Schizophrenia

SZ is a chronic mental disorder (Bhugra, 2005), which is typically characterized by cognitive problems, disintegration in perception of reality, auditory and/or visual hallucination, and a chronic course with lasting impairment (Heinrichs & Zakzanis, 1998). There is currently no standard clinical diagnostic test for SZ, and there has been considerable focus on identifying a biologically based marker using neuroimaging features which has shown some promise. We surveyed 101 peer-reviewed articles, which are presented in Table 1. Calhoun et al. (2004), Davatzikos, Shen, et al. (2005) and Yushkevich et al. (2005) are among the first studies to perform predictive analyses on SZ using MRI-based neuroimaging data (Table 1).

1 Structural MRI: By utilizing sMRI data, Davatzikos, Shen, et al. (2005) used voxel-based feature set and applied a high-dimensional nonlinear pattern classification approach to compute the degree of separation between SZ and HCs (HC). Using the leave-one-out CV (LOO-CV), the authors reported 81% classification accuracy (for gender-wise classification, 82% for women and 85% for men). Another study by Yushkevich et al. (2005) used SVM classifier and region-based feature sets to discriminate SZ patients with 72% accuracy. More recently, Koutsouleris et al. (2009) used sMRI and a principal component feature selection approach, where based on the overall predictive performance of the feature selection algorithm, an optimal number of principal components was identified to predict SZ. This study is particularly of importance as it reported to reliably predict different subcategories of SZ, with a three-class classification for SZ showing a maximal accuracy of 82%. Another large-scale study with a sample size of 256 case–control as well as a similar sized replication cohort predicted SZ based on sMRI-derived features with an accuracy of about 70% for both CV and replication study (Nieuwenhuis et al., 2012).

2 Functional MRI: More recently, a high number of studies used features from resting-state and task functional MRI (fMRI) for predictive modeling of SZ, and achieved promising outcomes.
| References                                      | Disorder | Modality | Number of subjects | Feature type | Classifier                  | Overall accuracy |
|-------------------------------------------------|----------|----------|--------------------|--------------|-----------------------------|------------------|
| Caprihan, Pearlson, and Calhoun (2008)          | SZ       | dMRI     | HC = 45, SZ = 45,  | Discriminant | Fisher’s LDC                | 80%              |
|                                                 |          |          | Total = 90         | PCA of FA maps |                             |                  |
| Caan et al. (2006)                              | SZ       | dMRI     | HC = 24, SZ = 34,  | FA map       | LDA                         | 75%              |
|                                                 |          |          | Total = 5          |              |                             |                  |
| Ardekani et al. (2011)                          | SZ       | dMRI     | HC = 50, SZ = 50,  | Voxels of FA| LDA                         | 96%              |
|                                                 |          |          | Total = 100        | and MA maps |                             |                  |
|                                                  |          |          |                    | reduced by   |                             |                  |
|                                                  |          |          |                    | PCA         |                             |                  |
| Honorio, Tomasi, Goldstein, Leung, and Samaras   | SZ       | fMRI     | HC = 15, SZ = 13,  | Mean activation | Majority vote of three | 96%              |
| (2012)                                          |          | (SM, AOD,| Total = 28         | of the largest | decision stumps            |                  |
|                                                 |          | WMT)     |                    | activation   |                             |                  |
| Hunorion, Tomasi, Goldstein, Leung, and Samaras  |          | fMRI     | HC = 91, SZ = 57,  | ICA spatial   | Projection pursuit         | 80–90%           |
| (2012)                                          |          | (AOD/Sternberg/| Total = 138       | maps         |                             |                  |
|                                                  |          | SM tasks) |                  |              |                             |                  |
| Demirci et al. (2008)                           | SZ       | fMRI     | HC = 51, SZ = 51,  | Voxels of left | LDA                         | 62%              |
|                                                 |          | (AX-CPT task) | Total = 102      | DLPFC in the |                             |                  |
|                                                 |          |          |                    | contrast map |                             |                  |
| Yoon et al. (2012)                              | SZ (first episode) | fMRI (SM task) | HC = 40, SZ = 32, | Sparse representation | Sparse representation-based classifier | 77%              |
|                                                 |          |          | BP = 40, Total = 104 | based variable selection |                             |                  |
| Koch et al. (2015)                              | SZ       | fMRI     | HC = 15, SZ = 19,  | Thresholded voxels in | SVM                  | 92%              |
|                                                 |          | (monetary incentive delay task) | Total = 34 | activation map by ANOVA tests |                             |                  |
| Cao, Duan, Lin, Calhoun, and Wang (2013)         | SZ       | fMRI     | HC = 116, SZ = 92, | Sparse representation | SVM                  | 75–91%           |
|                                                 |          | (SM task) and SNP | Total = 208     | based variable selection |                             |                  |
| CostaReda et al. (2011)                         | SZ/BP    | fMRI (verbal fluency task) | HC = 20, SZ = 16, | MVPA                  | SVM                  | 85%              |
|                                                 |          |          | BP = 40, Total = 53 | from the contrast map |                             |                  |
| Yoon et al. (2008)                              | SZ       | fMRI     | HC = 15, SZ = 19,  | Selected active voxels | MVPA                      | 59–72%           |
|                                                 |          | (visual task) | Total = 34       | from the contrast map  |                             |                  |
| Bleich-Cohen et al. (2014)                      | SZ with and without OCD | fMRI (WMT task) | HC = 20, SZ (with OCD) = 16, SZ (without OCD) = 17, Total = 53 | MVPA on GLM contrast values | SVM                   |                  |
| Castro, Gómez-Verdejo, Martínez-Ramón, Kiehl, and Calhoun (2014) |          |          |                    |                         |  |                  |
| Castro, Martínez-Ramón, Pearson, Sui, and Calhoun (2011) |         |          |                    | Multiple kernel learning | Recursive composite kernels | 95%              |
| Study                                      | Design  | Imaging Modality          | Group Comparison                           | Feature Extraction Method                                                                 | Classification Method                  | Accuracy     |
|-------------------------------------------|---------|---------------------------|--------------------------------------------|------------------------------------------------------------------------------------------|----------------------------------------|--------------|
| Calhoun, Maciejewski, Pearlson, and Kiehl (2008) | SZ/BP   | fMRI (AOD task)           | HC = 26, SZ = 21, BP = 14, Total = 61     | Distance to mean image for each group build using ICA spatial maps (DMN and temporal lobe) | Majority voting among 3 SVMs           | 87%          |
| Arribas et al. (2010)                     | SZ      | fMRI (AOD task)           | HC = 25, SZ = 21, BP = 14, Total = 60     | ICA spatial maps (DMN and temporal lobe)                                                 | Bayesian generalized Softmax perceptron | 82–90% (AUC) |
| W. Du et al. (2012)                       | SZ      | fMRI (AOD task) and rsfMRI| HC = 28, SZ = 28, Total = 56              | Kernel PCA on ICA spatial maps                                                            | Fisher’s LDC                             | 93–98%       |
| Çetin et al. (2015)                       | SZ      | fMRI (AOD task) and rsfMRI| HC = 28, SZ = 27, Total = 55              | FNC scores derived from ICA-based multinetwork fusion template                            | LDA and shaplet based classifier       | 72%          |
| Yang, Liu, Sui, Pearlson, and Calhoun (2010) | SZ      | fMRI (AOD task) and SNP   | HC = 20, SZ = 20, Total = 40              | Three types of features: Selected voxels in fMRI activation map, selected SNPs and ICA components | Majority voting among 3 SVMs           | 87%          |
| Yu et al. (2013)                          | SZ      | rsfMRI                    | HC = 25, SZ = 24, sibling, HC = 22, Total = 71 | FC among 116 regions in AAL atlas reduced by PCA                                        | SVM                                    | 62%          |
| Yu, Shen, Zeng, Ma, and Hu (2013)         | SZ/MDD  | rsfMRI                    | HC = 38, SZ = 32, MDD = 19, Total = 89    | FC among ROIs                                                                           | SVM                                    | 80.90%       |
| Watanabe, Kessler, Scott, Angstadt, and Sripada (2014) | SZ      | rsfMRI                    | HC = 74, SZ = 71, Total = 145             | FC among 347 nodes placed as a grid in the entire brain                                 | Fused lasso, GraphNet                  | 91%          |
| W. Guo et al. (2014)                      | SZ      | rsfMRI                    | HC = 46, unaffected sibling of SZ patients = 46, Total = 92 | fALFF values of the left ITG                                                             | SVM                                    | 75%          |
| Venkataraman, Whitford, Westin, Golland, and Kubicki (2012) | SZ      | rsfMRI                    | HC = 18, SZ = 18, Total = 36              | FC among 90 ROIs                                                                         | RF                                     | 75%          |
| Tang, Wang, Cao, and Tan (2012)           | SZ      | rsfMRI                    | HC = 22, SZ = 22, Total = 44              | FC among 90 ROIs reduced by PCA                                                          | SVM                                    | 93%          |
| Su, Wang, Shen, Feng, and Hu (2013)       | SZ      | rsfMRI                    | HC = 32, SZ = 32, Total = 64              | FC among 116 AAL regions                                                                 | SVM                                    | 83%          |
| Shen, Wang, Liu, and Hu (2010)            | SZ      | rsfMRI                    | HC = 20, SZ = 32, Total = 52              | Dimension-reduced FC (local linear)                                                       | C-means clustering                     | 86%          |
| Study                                      | Sample Size | Method | Sample Description | Features Used | Technique | Classification Accuracy |
|-------------------------------------------|-------------|--------|--------------------|---------------|-----------|------------------------|
| J. Kim et al. (2016)                      | SZ          | rsfMRI | HC = 50, SZ = 50,   | FC among 116  | DNN       | 86%                   |
|                                           |             |        | Total = 100        | regions in AAL|           |                        |
|                                           |             |        |                    | atlas         |           |                        |
| Kaufmann et al. (2015)                    | SZ          | rsfMRI | HC = 196, SZ = 71, | FC based on ICA| Regularized LDC | 75–84%     |
|                                           |             |        | Total = 267        | decomposition |           |                        |
| H. Cheng et al. (2015)                    | SZ          | rsfMRI | HC = 29, SZ = 19,  | Graph measures | SVM       | 80.00%                |
|                                           |             |        | Total = 48         | of FC         |           |                        |
| Fekete et al. (2013)                      | SZ          | rsfMRI | HC = 10, SZ = 8,   | Local and global | SVM       | 100%                  |
|                                           |             |        | Total = 18         | complex network|           |                        |
|                                           |             |        |                    | measures      |           |                        |
| Fan et al. (2011)                         | SZ          | rsfMRI | HC = 31, SZ = 31,  | FC patterns   | Ensemble of SVM classifiers | 85–87%     |
|                                           |             |        | Total = 62         |               |           |                        |
| Chyzhyk, Savio, and Graña (2015)          | SZ          | rsfMRI | HC = 74, SZ = 72,  | Pearson correlation | Ensemble of extreme learning| 80–91%     |
|                                           |             |        | Total = 146        | features derived | machines   |                        |
|                                           |             |        |                    | from ReHo, ALFF, |           |                        |
|                                           |             |        |                    | FALF and      |           |                        |
|                                           |             |        |                    | voxel-mirrored homotopic connectivity | |                        |
| Bassett, Nelson, Mueller, Camchong, and Lim (2012) | SZ          | rsfMRI | HC = 29, SZ = 29,  | Size of connected components in graphs build from | SVM       | 75%                   |
|                                           |             |        | Total = 58         | correlation among time-courses for 90 AAL regions |           |                        |
|                                           |             |        |                    |               |           |                        |
| Arbabshirani, Kiehl, Pearlson, and Calhoun (2013) | SZ          | rsfMRI | HC = 28, SZ = 28,  | FNC among ICA time-courses | SVM (best) | 96%                   |
|                                           |             |        | Total = 56         |               |           |                        |
| Anticevic et al. (2013)                   | SZ          | rsfMRI | HC = 90, SZ = 90,  | MVPA based on whole brain thalamic connectivity map | SVM       | 73.90%                |
|                                           |             |        | Total = 180        |               |           |                        |
| A. Anderson and Cohen (2013)              | SZ          | rsfMRI | HC = 74, SZ = 72,  | Graph metrics based on FNC computed from ICA | SVM       | 65%                   |
|                                           |             |        | Total = 146        |               |           |                        |
| T. Zhang and Davatzikos (2013)            | SZ          | sMRI   | HC = 79, SZ = 69,  | Voxels from five regions based on optimally discriminative VBM | SVM       | 71%                   |
|                                           |             |        | Total = 148        |               |           |                        |
| Zanetti et al. (2013)                     | SZ (first episode) | sMRI  | HC = 62, SZ = 62,  | Whole brain volumetric | SVM       | 73%                   |
|                                           |             |        | Total = 124        |               |           |                        |
| Study                                      | Diagnosis          | Imaging Modality | Sample Size | Voxel-wise/ROI-based Measurements | Analysis Method | Accuracy       |
|-------------------------------------------|--------------------|-----------------|-------------|----------------------------------|----------------|----------------|
| Takayanagi et al. (2011)                  | SZ (first episode) | sMRI            | HC = 40, SZ = 52, Total = 92 | Volume and mean CT of selected ROIs | DFA            | 80%            |
| Sun et al. (2009)                         | SZ and psychosis   | sMRI            | HC = 36, SZ = 36, Total = 72 | Cortical GMD | Sparse multinomial LRCr | 86%            |
| Schnack et al. (2014)                     | SZ/BP              | sMRI            | HC1 = 66, HC2 = 43, SZ1 = 66, SZ2 = 46, BP1 = 66, BP2 = 47, Total1 = 198, Total2 = 136 | Voxel-wise GM maps | SVM            | 67–90%         |
| Radulescu et al. (2014)                   | SZ                 | sMRI            | HC = 24, SZ = 27, Total = 51 | Texture and volumetric measures | LDA            | 65.0–72.7%     |
| Pina-Camacho et al. (2015)                | SZ                 | sMRI            | HC = 42, SSD = 36, non-SSD = 45, Total = 123 | Clinical, neuropsychological, biochemical and volumetric measures | SVM            | 81.0–99.0%     |
| Pardo et al. (2006)                       | SZ/BP              | sMRI            | HC = 8, SZ = 10, BP = 10, Total = 28 | Volume of 23 ROIs along with 22 neuropsychological test scores | LDA            | 96%            |
| Ota et al. (2012)                         | SZ                 | sMRI            | HC = 105, HC2 = 23, SZ1 = 38, SZ2 = 23, Total = 189 | GM and CSF volumetric measures of ROIs | LDA            | 70–76%         |
| Nieuwenhuis et al. (2012)                 | SZ                 | sMRI            | HC1 = 111, HC2 = 122, SZ1 = 128, SZ2 = 155, Total1 = 239, Total2 = 277 | GMs based VBM of top 10% voxels | SVM            | 71%            |
| Nakamura et al. (2004)                    | SZ                 | sMRI            | HC = 47, SZ = 57, Total = 104 | Volume of several ROIs in the brain | LDA            | 78–86%         |
| Koutsouleris et al. (2015)                | SZ/MD              | sMRI            | MD = 104, SZ = 158, Total = 262 | GM maps of regional analysis of brain volumes in normalized space (RAVENS) | SVM-RFE        | 76%            |
| Kawasaki et al. (2007)                    | SZ                 | sMRI            | HC = 46, SZ = 46, Total = 92 | The mean expression of Eigen image derived from VBM | Simple thresholding | 80–90%         |
| Kasparek et al. (2011)                    | SZ (first-episode) | sMRI            | HC = 39, SZ = 39, Total = 78 | Whole brain voxel intensity values | Maximum-uncertainty LDA | 72%            |
| Study                                      | Group (onset) | Imaging Modality | Sample Size | Technique/Measurements                                                                 | Classification/Algorithm | Accuracy  |
|-------------------------------------------|---------------|-----------------|-------------|----------------------------------------------------------------------------------------|--------------------------|-----------|
| Karageorgiou et al. (2011)                | SZ (recent onset) | sMRI            | HC = 47, SZ = 28, Total = 75 | Volumetric measurements of 95 ROIs                                                   | LDA                      | 72%       |
| Janousova, Schwarz, and Kasparek (2015)   | SZ             | sMRI            | HC = 49, SZ = 49 | MR intensities, GMDs, and deformation based morphometry                               | Combination of mLDA, centroid method, and the average linkage | 81.60%    |
| Ivabuchi, Liddle, and Palaniyappan (2013) | SZ             | sMRI            | HC = 20, SZ = 19, Total = 39 | GM and WM maps                                                                        | SVM                      | 66.6–77%  |
| Ingalihalikar et al. (2012)               | SZ (identifying subtypes) | sMRI            | HC = 29, SZ = 23, Total = 52 | Multi-edge graphs build from structural connectivity networks with 78 ROIs          | Spectral clustering       | 78%       |
| Greenstein, Weisinger, Mailey, Clasen, and Gogtay (2012) | SZ (childhood onset) | sMRI            | HC = 99, SZ = 98, Total = 197 | CT                                      | RF                      | 74%       |
| Gould et al. (2014)                       | SZ (cognitive deficit and cognitive spared) | sMRI            | HC = 163, SZ = 208, SZA = 41, Total = 412 | Whole brain VBM                                                                       | SVM                      | 56–72%    |
| Fan, Shen, Gur, Gur, and Davatzikos (2007) | SZ             | sMRI            | HC1 = 38, HC2 = 41, SZ1 = 23, SZ2 = 46, Total 1 = 61, Total 2 = 87 | Volumetric measurements based on deformation-based morphometry | SVM                      | 91%       |
| Fan, Shen, and Davatzikos (2005)          | SZ             | sMRI            | HC = 38, SZ = 23, Total = 61 | Volumetric measures of all WM, GM, and CSF                                           | SVM-RFE                  | 92%       |
| Davatzikos et al. (2005)                  | SZ             | sMRI            | HC = 79, SZ = 69, Total = 148 | Whole brain volumetric measurements                                                   | Nonlinear classifier (n.s.) | 81%       |
| Csemansky et al. (2004)                   | SZ             | sMRI            | HC = 65, SZ = 52, Total = 117 | Hippocampal and thalamic shape eigenvectors                                           | DFA                      | 79%       |
| Castellani et al. (2012)                  | SZ             | sMRI            | HC = 54, SZ = 54, Total = 108 | Visual words extracted from DLPFC by SIFT and clustered by k-means                    | SVM with local kernel     | 66–75%    |
| Bansal et al. (2012)                      | SZ             | sMRI            | HC = 40, SZ = 65, Total = 105 | Semi-supervised (hierarchical clustering)                                            |                          | 94.00%    |

(Continued)
| Study                        | Group | Imaging Techniques | Measures                                      | Classification Method | Accuracy |
|------------------------------|-------|--------------------|-----------------------------------------------|-----------------------|----------|
| Ota et al. (2013)            | SZ/ MDD | sMRI and dMRI      | Volume and FA of insula, thalamus, ACC, ventricles and corpus callosum | LDA                  | 72–88%   |
| Sui et al. (2013)            | SZ    | sMRI, rsfMRI, and dMRI | GMDs from structural, FA from DTI and ALFF from fMRI | SVM                  | 79%      |
| Yushkevich et al. (2005)     | SZ    | sMRI               | Normalized regional volumes                   | N.S.                 | 71%      |
| Davatzikos, Shen, et al. (2005) | SZ    | sMRI               | GM, WM, and ventricular CSF volumes           | N.S.                 | 81%      |
| Shi et al. (2007)            | SZ    | sMRI               | ReHo                                          | Pseudo-Fisher LDA    | 80%      |
| Pohl and Sabuncu (2009)       | SZ    | sMRI               | Structure-specific transformation parameters   | SVM                  | 90%      |
| Rath et al. (2010)           | SZ    | dMRI               | Diffusion measures                            | Multiple classifiers | 85.50%   |
| Mueller, Wang, Pan, Holt, and Liu (2015) | SZ    | rsfMRI             | Regional specialization of caudate nucleus    | SVM                  | 74%      |
| Pouyan and Shahamat (2015)   | SZ    | rsfMRI             | Independent components                        | Majority vote method using multiple classifiers | 100%     |
| Lee et al. (2018)            | SZ    | sMRI and dMRI      | Structural measures (volume and/or FA)        | RF and SVM           | 87.6% - 89.5% |
| Pergola et al. (2017)        | SZ    | sMRI               | Thalamic GMV                                  | RF                   | 81% - 75% |
| Y. Liu et al. (2018)         | SZ    | rsfMRI             | Adolescent-onset schizophrenia (AOS) = 48, HC = 31, Total = 79 | Cohe-ReHo | 89.87%   |
| Winterburn et al. (2017)     | SZ    | sMRI               | Dataset 1: SZ = 88, HC = 103, Dataset 2: SZ = 91, HC = 67, Tissue density estimates and CT | LR, SVM and LDA      | 73.5% (best) |
| Study Reference                  | Group| Imaging Modality | Dataset Details | Analysis Details | Methodology | Accuracy |
|---------------------------------|------|------------------|-----------------|------------------|-------------|----------|
| Huang et al. (2018)             | SZ   | rsfMRI           | SZ = 50, HC = 50, Total = 449 | FC between 116 ROIs from AAL atlas | SVM         | 81.30%   |
| J. Shao et al. (2017)           | SZ   | dMRI             | SZ = 21, MDD = 25, HC = 25, Total = 71 | Structural connectivity from ROIs | Multiple classifiers | 80.4% (best) |
| Liu et al. (2017)               | SZ   | sMRI and dMRI    | SZ = 62, HC = 33, Total = 95 | GM and WM features | MFMK-SVM | 91.28%   |
| Vyškovský, Schwarz, Janoušová, and Kasparek (2016) | SZ   | sMRI             | SZ = 52, HC = 52, Total = 104 | GMD | MLP and SVM | 68% (best) |
| Ulloa, Plis, and Calhoun (2018) | SZ   | sMRI and fMRI    | SZ = 135, HC = 169, Total = 304 | GM map and ALFF maps | Multi-layer perceptron model | 85% (ROC) |
| S. Liang et al. (2018)          | SZ   | sMRI and dMRI    | Dataset 1: First-episode schizophrenia (FES) = 98, HC = 106; Dataset 2: FES = 54, HC = 48, Total = 306 | Regional GMV, CT, gyrification, FA, and MD | GBDT | 75.05% |
| P Mikolas et al. (2016)         | SZ   | rsfMRI           | Schizophrenia-spectrum disorder (FES) = 63, HC = 63, Total = 126 | Seed-based FC maps | SVM | 73.00% |
| de Pierrefeu et al. (2018)      | SZ   | sMRI             | SZ = 276, HC = 330, Total = 606 | Multiple structural measures (GM VBM, vertex-based CT) | Enet-TV penalty | 72% |
| Moghimi, Lim, and Netoff (2018) | SZ   | rsfMRI           | Chronic SZ = 52, first episode = 30, HC = 88, Total = 170 | Graph-theoretic measures | SVM | 80% |
| Janousova, Montana, Kasparek, and Schwarz (2016) | SZ   | sMRI             | First-episode SZ = 52, HC = 52, Total = 104 | GM volumes and local deformation from DBM | Multiple classifiers | 83.7% (best) |
| Salvador et al. (2017)          | SZ   | sMRI             | SZ = 128, BP = 128, HC = 127, Total = 383 | sMRI features including GM and WM VBM, vertex-based CT and volume, ROI volumetric measures and WBM maps | Multiple classifiers | 75% (best) |
| Study                        | Group | Imaging | Sample Size | Methodology | Features                  | Classifier | Accuracy |
|------------------------------|-------|---------|-------------|-------------|---------------------------|------------|----------|
| Kottaram et al. (2018)       | SZ    | rsfMRI  | SZ = 41, HC = 41, Total = 82 | Static and dynamic FC measures | SVM         | 91%      |
| Dagnew, Squarcina, Rivolta, Brambilla, and Sassi (2017) | SZ    | sMRI    | SZ = 20, HC = 20, Total = 40 | Features characterizing brain anatomy | SVM and KNN | 74%      |
| Lu et al. (2016)             | SZ    | sMRI    | SZ = 42, HC = 42, Total = 84 | GMV and WMV between | SVM         | 88.40%   |
| Tas et al. (2018)            | SZ    | sMRI    | SZ = 23, schizo-obsessive = 23, Total = 46 | Voxel of interest values | SVM         | 78.26%   |
| W. Cheng et al. (2015)       | SZ    | rsfMRI  | SZ = 415, HC = 405, Total = 820 | Resting-state thalamocortical FC | SVM         | 73.5% – 80.92% |
| Liu et al. (2017)            | SZ    | sMRI    | SZ = 38, HC = 38, Total = 76 | CT | SVM-RBF | 88.72% |
| Pavol Mikolas et al. (2018)  | SZ    | dMRI    | First episode SZ = 77, HC = 77, Total = 154 | FA | SVM | 62.34% |
| Q. Zhu, Huang, and Xu (2018) | SZ    | rsfMRI  | SZ = 24, HC = 21, Total = 45 | FC measures | KDA | 95.56% |
| B. Cao et al. (2018)         | SZ    | rsfMRI  | SZ = 43, HC = 29, Total = 72 | FC measures | SVM | 78.60% |
| Gheiratmand et al. (2017)    | SZ    | Task fMRI (AOD) | SZ = 184, HC = 196, Total = 380 | Whole-brain FC measures | Multiple classifiers | 74% |
| Chin, You, Meng, Zhou, and Sim (2018) | SZ    | sMRI    | SZ = 141, HC = 71, Total = 212 | GMD | SVM | 86.6% (recognition) |
| Guo, Palaniyappan, Liddle, and Feng (2016) | SZ    | sMRI    | SZ = 98, HC = 83, Total = 181 | CT | SVM | 96.30% |
| Cabral et al. (2016)         | SZ    | rsfMRI and sMRI | SZ = 74, HC = 71, Total = 145 | GM volume and FC measures | MVPA | 75% |
| Rozycki et al. (2017)        | SZ    | sMRI    | SZ = 440, HC = 501, Total = 941 | Regional volumes, voxel-wise measures, and complex distributed patterns | SVM | 76% |
| Dwyer et al. (2018)          | SZ    | sMRI    | SZ = 71, HC = 74, Total = 145 | GMV | SVM | 68.5–73% |
| References                        | Disorder        | Modality     | Number of subjects | Feature type | Classifier | Overall accuracy |
|----------------------------------|-----------------|--------------|--------------------|--------------|------------|-----------------|
| Rubin-Falcone et al. (2018)      | BP/MDD          | sMRI         | BP = 26, MDD = 26, Total = 52 | GM           | SVM        | 75%             |
| Deng et al. (2018)               | BP/MDD          | DTI          | BP = 31, MDD = 36, Total = 67 | FA           | SVM        | 68.30%          |
| Gao et al. (2017)                | BP/MDD          | rsfMRI       | BP = 37, MDD = 36, Total = 73 | Spatial independent components | SVM        | 93.00%          |
| Jing et al. (2017)               | cMDD/rMDD       | rsfMRI       | cMDD = 19, rMDD = 19, HC = 19, Total = 57 | Hurst exponent | SVM        | 87.0% (cMDD vs. HC), 84.0% (rMDD vs. HC), 89.0% (cMDD vs. rMDD) |
| Yoshida et al. (2017)            | MDD/BP          | rsfMRI       | MDD = 58, HC = 65, Total = 123 | FC           | PLS        | 80.00%          |
| M. Li et al. (2017)              | MDD/BP          | rsfMRI       | BP = 22, MDD = 22, Total = 44 | Degree centrality | SVM        | 86.00%          |
| Zhong et al. (2017)              | MDD             | rsfMRI       | MDD1 = 29, HC1 = 33, MDD2 = 46, HC2 = 57, Total1 = 62, Total2 = 103 | FC           | SVM        | 91.9% (first sample), 86.4% (second sample) |
| Wang, Ren, and Zhang (2017)      | MDD             | rsfMRI       | MDD = 31, HC = 29, Total = 60 | FC           | SVM        | 95.00%          |
| Schnyder, Clasen, Gonzalez, and Beevers (2017) | MDD             | DTI          | MDD = 25, HC = 25, Total = 50 | FA           | SVM        | 74.00%          |
| Sundermann et al. (2017)         | MDD             | rsfMRI       | MDD = 180, HC = 180, Total= | FC           | SVM        | 45.0% – 56.1%   |
| Bhuamik et al. (2017)            | MDD             | rsfMRI       | MDD = 38, HC = 29, Total= | FC           | SVM        | 76.10%          |
| He et al. (2017)                 | MDD/BP          | rsfMRI/sMRI  | BP = 13, MDD = 40, HC = 33, Total= | FNC, GM       | SVM        | 91.3% (three groups), 99.0% (BD vs. MDD) |
| Bürger et al. (2017)             | MDD/BP          | Task fMRI    | BP = 36, MDD = 36, HC = 36, Total= | Contrast maps | SVM, GPC     | 72.00%          |
| Hilbert, Lueken, Muehlan, and Beesdo-Baum (2017) | GAD/MDD          | sMRI         | GAD = 19, MDD = 14, HC = 24, Total= | GM           | SVM        | 58.7% (GAD and MDD vs. HC), 68.1% (GAD vs. MDD) |
| Drysdale et al. (2017)           | MDD (biotypes)  | rsfMRI       | MDD = 333(4 biotypes), HC = 378, Total= | FC           | SVM        | 89.20%          |
| Sankar et al. (2016)             | MDD             | sMRI         | MDD = 23, HC = 20, Total= | GM, WM       | SVM        | 70.00%          |
| Frangou et al. (2017)            | MDD/BP          | Task fMRI    | BP = 30, MDD = 30, Total= | Contrast maps | GPC        | 73.10%          |
| Ramasubbu et al. (2016)          | MDD             | rsfMRI       | MDD = 15, HC = 19, Total= | Spatial independent components | SVM        | 66.00%          |
| W. Yang et al. (2016)            | MDD             | Task fMRI    | MDD = 16, HC = 16, Total= | Contrast maps | SVM        | 75.00%          |
| Rive et al. (2016)               | MDD/BP          | sMRI/rsfMRI  | MDr = 23, BPr = 26, MDD = 22, BNI = 10, Total = 81 | GM, spatial independent components | GPC        | 69.10%          |
| N.-F. Jie et al. (2015)          | MDD/BP          | sMRI         | BP = 21, MDD = 25, Total = 46 | GM, fALFF    | SVM        | 92.10%          |
| Foland-Ross et al. (2015)        | MDD             | sMRI         | MDD = 18, HC = 15, Total = 33 | CT           | SVM        | 69.70%          |
| Sacchet, Livermore, Iglesias, Glover, and Gotlib (2015) | MDD/BP          | sMRI         | BP = 40, MDD = 57, HC = 61, Total = 158 | GM           | SVM        | 59.5% (BP vs. MDD), 62.8% (MDD vs. HC) |

(Continues)
| Study                          | Type       | Group I | Group II | Total | Graph metrics | Methodology       | Accuracy   |
|-------------------------------|------------|---------|----------|-------|---------------|-------------------|------------|
| Sacchet, Prasad, Foland-Ross, Thompson, and Gotlib (2015) | MDD, DTI  | MDD = 14, HC = 18, Total = 32 | Graph metric of WM connectivity | SVM | 71.90% |
| (João R Sato et al. 2015)     | MDD        | MDD = 25, HC = 21, Total = 46 | Contrast maps | LDA | 78.30% |
| Johnston, Steele, Tolomeo, Christmas, and Matthews (2015) | MDD        | MDD = 20, HC = 21, Total = 41 | GM | SVM | 85.00% |
| Johnston et al. (2015)        | MDD        | MDD = 19, HC = 21, Total = 40 | Contrast maps | SVM | 76.00% |
| Koutsouleris et al. (2015)    | MDD        | MDD = 104, SZ = 158, Total = 262 | GM | SVM | (best) |
| (Shimizu et al. 2015))        | MDD        | MDD = 31, HC = 31, Total = 62 | Contrast maps | gLASSO, SVM | 95.0% (best) |
| Fung et al. (2015)            | MDD/BP     | BP = 16, MDD = 19, Total = 35 | CT and surface area | SVM | 74.30% |
| Rosa et al. (2015)            | MDD        | MDD = 19, HC = 19, Total = 38 | FC | SVM | 85.00% |
| Patel et al. (2015)           | MDD        | MDD = 33, HC = 35, Total = 68 | rsfMRI, sMRI, DTI | Decision tree | 87.30% |
| Redlich et al. (2014)         | MDD/BP     | BP = 58, MDD = 58, Total = 116 | GM | GPC | 79.30% |
| L. Cao et al. (2014)          | MDD        | MDD = 39, HC = 37, Total = 76 | FC | SVM | 84.00% |
| MacMaster, Carrey, Langevin, Jaworska, and Crawford (2014) | MDD/BP     | BP = 14, MDD = 32, Total = 43 | GM | LDA | 81.00% |
| L. L. Zeng, Shen, Liu, and Hu (2014) | MDD, rsfMRI | MDD = 24, HC = 29, Total = 53 | FC | MMC | 92.5% (clustering), 92.5% (classification) |
| Rondina et al. (2014)         | MDD        | MDD = 30, HC = 30, Total = 60 | Voxel intensity | SVM | 72.00% |
| H. Guo, Su, et al. (2014)     | MDD        | MDD = 36, HC = 27, Total = 63 | FC | NN | 90.50% |
| (Serpa et al. 2014))          | MDD/BP     | BP = 23, MDD = 19, HC = 38, Total = 80 | GM, WM, and ventricular RAVENS maps | SVM | 54.8% (BD vs. MDD), 59.6% (MDD vs. HC) |
| Habes et al. (2013)           | MDD        | MDD = 9, HC = 9, Total = 18 | Contrast maps | LDA | 72.20% |
| Wei et al. (2013)              | MDD        | MDD = 20, HC = 20, Total = 40 | Spatial independent components | SVM | 90.00% |
| Grotegerd et al. (2014)       | MDD/BP     | BP = 22, MDD = 22, Total = 44 | Contrast maps | GPC | 79.60% |
| Yu, Shen, Zeng, et al. (2013) | MDD        | MDD = 19, SZ = 32, Total = 51 | FC | SVM | 80.90% |
| Modinos et al. (2013)         | MDD        | MDD = 17, HC = 17, Total = 34 | Contrast maps | SVM | 77.00% |
| Ma, Li, Yu, He, and Li (2013) | MDD        | MDD = 19, HC = 18, Total = 37 | ReHo | LDA | 91.90% |
| Grotegerd et al. (2013)       | MDD/BP     | BP = 10, BP = 10, HC = 10, Total = 30 | Contrast maps | SVM, GPC | 90.00% |
| Fang et al. (2012)            | MDD        | MDD = 22, HC = 26, Total = 48 | Anatomical connectivity | SVM | 91.70% |
| Study                                      | Region       | Imaging Technique | MDD | HC | Total | Feature | Voxel-based Feature (GM) | Method | Accuracy |
|--------------------------------------------|--------------|-------------------|-----|----|-------|---------|--------------------------|--------|----------|
| Mwangi, Ebmeier, Matthews, and Douglas Steele (2012) | MDD          | sMRI              | MDD = 62, HC = 62, Total = 124 | Voxel-based feature (GM) | RVM | 90.30% |
| Lord, Hor, Breakspear, and Walter (2012)   | MDD          | rsfMRI            | MDD = 22, HC = 22, Total = 44 | FC | SVM | 99.00% |
| F. Liu et al. (2012)                       | TRD/TSD      | sMRI              | TRD = 18, TSD = 17, HC = 17, Total = 52 | GM, WM | SVM | 82.90% |
| L.-L. Zeng et al. (2012)                   | MDD          | rsfMRI            | MDD = 24, HC = 29, Total = 53 | FC | SVM | 94.30% |
| Mourão-Miranda et al. (2011)               | MDD          | Task fMRI         | MDD = 19, HC = 19, Total = 38 | Contrast maps | SVM | 52.0% (true positive) |
| Hahn et al. (2011)                         | MDD          | Task fMRI         | MDD = 30, HC = 30, Total = 60 | Contrast maps | GPC | 83.00% |
| Nouretdinov et al. (2011)                  | MDD          | Task fMRI         | MDD = 19, HC = 19, Total = 38 | Contrast maps | SVM | 76.30% |
| Costafreda, Chu, Ashburner, and Fu (2009)  | MDD          | sMRI              | MDD = 37, HC = 37, Total = 74 | GM | SVM | 67.60% |
| Fu et al. (2008)                           | MDD          | Task fMRI         | MDD = 19, HC = 19, Total = 38 | Contrast maps | SVM | 86.00% |
| Matsuo et al. (2018)                       | MDD/BP       | sMRI              | MDD = 59, BP = 158, HC = 777, Total = 1,531 | GMV | SVM | 88.1% (best) |
| Geng, Xu, Liu, and Shi (2018)              | MDD          | rsfMRI            | MDD = 24, HC = 24, Total = 48 | FC and EC | Linear and nonlinear SVM, KNN, LR | 91.70% |
| Rubin-Falcone et al. (2018)                | MDD/BP       | sMRI              | MDD = 26, BP = 26, Total = 52 | GMV | SVM | 75% |
| Nunes et al. (2018)                        | BP           | sMRI              | BP = 85, HC = 2,167, Total = 3,020 | Regional CT, surface area and subcortical volume | SVM | 81.10% |
| J. Yang et al. (2018)                      | MDD          | dMRI and sMRI     | Test dataset: MDD = 147, HC = 52; validation dataset: MDD = 83, HC = 25, Total = 307 | Multiple structural measures | Penalized LR, RF and SVM | 75.10% |
| References                                      | Disorder | Modality       | Number of subjects | Feature type                                      | Classifier          | Overall accuracy |
|------------------------------------------------|----------|----------------|--------------------|--------------------------------------------------|---------------------|------------------|
| Ingalhalikar, Parker, Bloy, Roberts, and Verma (2011) | ASD      | dMRI           | TDC = 30, ASD = 45, Total = 75 | FA and MD of selected ROIs | SVM                 | 80%              |
| Just, Cherkassky, Buchweitz, Keller, and Mitchell (2014) | ASD      | fMRI (social interaction task) | HC = 17, TDC = 17, Total = 34 | Activation of selected voxels processed by factor analysis | GNB                 | 97%              |
| Murdough et al. (2012)                                | ASD      | fMRI (two language tasks and a theory-of-mind task) | TD = 14, ASD = 13, Total = 30 | AG, MPFC and PCC based FC maps                     | LR                  | 96.00%           |
| Deshpande, Libero, Sreenivasan, Deshpande, and Kana (2013) | ASD      | Task fMRI and DMRI | TDC = 15, ASD = 15, Total = 30 | Causal connectivity weights, FC values and FA values | SVM                 | 95.90%           |
| Uddin et al. (2013)                                  | ASD      | rsfMRI         | TDC = 20, ASD = 20, Total = 40 | ICA components of rsfMRI                          | LR                  | 78.00%           |
| Plitt et al. (2015)                                  | ASD      | rsfMRI         | TD1 = 59, TD2 = 89, ASD1 = 59, ASD2 = 89, Total = 296 | FC among ROIs | LR and SVM (best) | 76.70%           |
| Iidaka (2015)                                       | ASD      | rsfMRI         | TDC = 328, ASD = 312, Total = 640 | FC among 90 ROIs | Probabilistic neural network | 90%              |
| C. P. Chen et al. (2015)                             | ASD      | rsfMRI         | TDC = 126, ASD = 126, Total = 252 | FC among 220 ROIs | RF                  | 91%              |
| J. S. Anderson et al. (2011)                         | ASD      | rsfMRI         | TD = 40, ASD = 40, Total = 80 | FC among ROIs | Thresholding       | 79.00%           |
| Wee, Wang, Shi, Yap, and Shen (2014)                 | ASD      | sMRI           | HC = 59, ASD = 58, Total = 117 | Thickness and volumetric of ROIs along with interregional features | Multi-kernel SVM | 96.30%           |
| Uddin et al. (2011)                                  | ASD      | sMRI           | TD = 24, ASD = 24, Total = 48 | Voxel-wise GM and WM maps | SVM                 | 92.00%           |
| Segovia et al. (2014)                                | ASD      | sMRI           | HC = 40, ASD = 52, ASD-sib = 40 | GM volume map | SVM                 | 80.0–85.0%       |
| Jiao et al. (2010)                                   | ASD      | sMRI           | HC = 16, ASD = 22, Total = 38 | Regional thickness measurements extracted from SBM | Logistic model trees | 87%              |
| Gori et al. (2015)                                   | ASD      | sMRI           | HC = 20, ASD = 21, Total = 41 | Morphometric features of selected ROIs            | SVM                 | 74% (AUC)        |
| Ecker, Rocha-Rego, et al. (2010)                     | ASD      | sMRI           | HC = 22, ASD = 22, Total = 44 | GM and WM maps | SVM                 | 77%              |
| Ecker et al. (2010)                                  | ASD      | sMRI           | HC = 20, ASD = 20, Total=40 | Volumetric and geometric features of selected cortical locations | SVM                 | 85%              |
| Calderoni et al. (2012)                              | ASD      | rsfMRI         | TDC = 38, ASD = 30, Total = 76 | Gray matter maps | SVM                 | 80% (AUC)        |
| Akshoomoff et al. (2004)                             | ASD      | sMRI           | TDC = 15, ASD = 52, Total = 67 | Volumetric measures and cerebellar vermis area    | DFA                 | 92.3–95.8%       |
| Study                          | Group | Imaging Modalities | Sample Size | Metrics/Methods | Classification Model | Accuracy |
|-------------------------------|-------|--------------------|-------------|-----------------|----------------------|----------|
| Libero, DeRamus, Lahti, Deshpande, and Kana (2015) | ASD | sMRI, dMRI and MRS | TD = 18, ASD = 19, Total = 37 | Cortical thickness, FA and neurochemical concentration | Decision tree | 91.90% |
| Zhou, Yu, and Duong (2014) | ASD | sMRI, rsfMRI | TDC = 153, ASD = 127, Total = 280 | Volume of selected subcortical regions, fALFF, number of voxels and Z-values of selected regions and global VMHC voxel Number | Random tree classifier | 70% |
| F. Zhang et al. (2018) | ASD | dMRI | ASD = 70, TD = 79, Total = 149 | FA and MD measures | SVM | 78.33% |
| Bernas, Aldenkamp, and Zinger (2018) | ASD | rsfMRI | Dataset 1: ASD = 12, HC = 12; Dataset 2: ASD = 12, HC = 18, Total = 54 | Time of in-phase coherence | LDA and SVM | 86.70% |
| Chanel et al. (2016) | ASD | Task fMRI (social stimuli) | ASD = 15, HC = 14, Total = 29 | Beta map voxels | SVM-RFE | 69–92.3% |
| Dodonova, Belyaev, Tkachev, Petrov, and Zhukov (2016) | ASD | dMRI | ASD = 51, TD = 43, Total = 94 | Graph spectral distributions | EMD-based kernel | 71% (AUC) |
| Tejwani, Liska, You, Reinen, and Das (2017) | ASD | rsfMRI | ASD = 147, HC = 146, Total = 293 | Dynamic FC variability | Naive Bayes, RF, SVM and multilayer perceptron algorithm | 65% |
| Katuwal, Baum, and Michael (2018) | ASD | sMRI | ASD = 15, non-ASD = 18, Total = 33 | Multiple brain morphometric features | RF | 95% (AUC) |
| Bhaumik, Pradhan, Das, and Bhaumik (2018) | ASD | rsfMRI | ASD = 145, HC = 159, Total = 372 | FC measures from ROIs | MVPA | 70% (best) |
| Fredo, Jahedi, Reiter, and Müller (2018) | ASD | rsfMRI | Training: ASD = 160, HC = 160; validation: ASD = 40, HC = 40, Total = 400 | FC measures from ROIs | RF | 65% |
| Zhao, Zhang, Rekik, An, and Shen (2018) | ASD | rsfMRI | ASD = 54, HC = 46, Total = 100 | FC measures | SVM | 81% |
| Soussia and Rekik (2018) | ASD | sMRI | ASD = 155, HC = 186, Total = 341 | Network-based measures | Supervised ensemble classifier | 61% (best) |
| Retico, Gori, Giuliano, Muratori, and Calderoni (2016) | ASD | sMRI | ASD = 41, HC = 40, Total = 81 | Surface-based measures | SVM | 74% (AUC, best) |
| Ghiassian, Greiner, Jin, and Brown (2016) | ASD | rsfMRI & sMRI | ADHD dataset: ADHD = 279, HC = 490; ASD dataset: ASD = 77, HC = 94, Total = 940 | sMRI and fMRI measures | HOG- MHPC | 65.00% |
| Dekhil et al. (2018) | ASD | rsfMRI | ASD = 123, TD = 160, Total = 283 | Higher level fMRI features from SAEs | Probabilistic SVM | 92% (best) |
| L. Wang, Wee, Tang, Yap, and Shen (2016) | ASD | sMRI | ASD = 54, HC = 57, Total = 111 | GM and WM features | SVM | 75.40% |

(Continues)
(a) **Task-based**: Studies using features from task-based fMRI paradigms include experiments with verbal fluency, working memory and auditory oddball (Castro et al., 2011; Costafreda et al., 2011; Honorio et al., 2012). One of the first relatively large-scale study (i.e., sample size >150) classified SZ using three different task-based fMRI data (i.e., auditory oddball, Sternberg item recognition and sensorimotor tasks) of 155 participants from two sites (Demirci, Clark, Magnotta, et al., 2008). The authors applied a projection pursuit algorithm on ICA spatial maps, and achieved classification accuracies ranging between 80 and 90%, with sensorimotor task providing the best performance. Further, based on regions with greater synchrony estimated from synchronous hemodynamic independent maps of auditory cortex as features, Calhoun and colleagues used a within-participant subtractive comparison to discriminate SZ from HC with 97% initial accuracy and 94% accuracy after a retest validation using a new subjects scanned at a different site (Calhoun et al., 2004). Another study focusing on three-class differential diagnosis of SZ, BP and HC individuals reported that verbal fluency led to a reliable diagnostic specificity for SZ with 92% accuracy (Costafreda et al., 2011). Many psychotic disorders, such as SZ, schizoaffective and BP disorders may share substantial number of overlapping symptoms, risk genes, brain dysfunctions, and treatment response. Thus, it becomes very challenging to clinically differentiate these patients based on traditional diagnostic approaches. A few recent studies have investigated the heterogeneity associated with SZ and schizoaffective disorder, as well as BP disorder with psychosis (Cardno & Owen, 2014; Cosgrove & Suppes, 2013; Pearson, Clementz, Sweeney, Keshavan, & Tamminga, 2016). Clementz and colleagues proposed a “biotypes” based approach where they identified three neurobiologically unique biologically defined psychosis categories and shown that biotypes did not follow a straightforward disease severity continuum, with heritable properties in unaffected first-degree relatives (Clementz et al., 2015).

(b) **Resting-state**: The rsfMRI studies for SZ prediction included a variety of classifiers, such as, SVM, fused lasso, GraphNet, RF, C-means clustering, regularized LDC, and ensembles of SVM classifiers, as presented in Table 1. Overall, the sample size was relatively large across these studies, with classification accuracies ranging from 62% to 100%, although the study with 100% accuracy rate had a very small sample size (20 participants) and therefore, the results might not be generalizable across other studies (Pouyan & Shahamat, 2015).

3 **Diffusion MRI**: The dMRI studies surveyed in this review reported accuracies ranging from 62% to 96%, with classifiers such as SVM, LDA, Fisher’s LDC or combination of multiple classifiers (Ardekani et al., 2011; Caan et al., 2006; Caprihan et al., 2008; Mikolas et al., 2018; Rathi et al., 2010; J. Shao et al., 2017). Features from these studies included fractional anisotropy (FA) maps and structural connectivity from ROIs.

4 **Multimodal**: Recently, by using connectivity measures from multimodal dMRI and sMRI data, Zhu, Shen, Jiang, and Liu (2014) predicted SZ patients and achieved a perfect accuracy (i.e., 100%). However, given the small sample size (i.e., HC = 10 and SZ = 10) and the choice of CV (i.e., LOO-CV), the framework might have introduced classification bias with lack of generalizability with independent study sample.
| References                     | Disorder                  | Modality                  | Number of subjects | Feature type | Classifier        | Overall accuracy |
|-------------------------------|---------------------------|---------------------------|--------------------|--------------|-------------------|------------------|
| Hart et al. (2014)            | ADHD                      | fMRI (stop task)         | HC = 30, ADHD = 30, Total = 60 | Whole brain GLM coefficient map | GPC               | 77%              |
| Park, Kim, Seo, Lee, and Park (2016) | ADHD                      | fMRI (six task)          | ADHD-IA = 13, ADHD-C = 21, Total = 34 | Network measures based on FC values | SVM               | 91.20%           |
| Hart et al. (2014)            | ADHD                      | fMRI (temporal discrimination task) | HC = 20, ADHD = 20, Total = 40 | Brain activation map | GPC               | 75.00%           |
| (Zou & Hastie, 2005)          | ADHD                      | rsfMRI                    | HC = 12, ADHD = 12, Total = 24 | ReHo maps | PCA-based fisher discriminative analysis | 85.00%           |
| Wang, Jiao, Tang, Wang, and Lu (2013) | ADHD                      | rsfMRI                    | HC = 23, ADHD = 23, Total = 46 | ReHo maps | SVM               | 80.00%           |
| Dey, Rao, and Shah (2014)     | ADHD                      | rsfMRI                    | HC = 429, ADHD-I = 98, ADHD-C = 141, Total = 668 | FFT and different variation of PCA on the BOLD signals along with phenotypic measures | SVM               | 68.86–76%        |
| Sato, Hoexter, Fujita, and Rohde (2012) | ADHD                      | rsfMRI                    | HC = 546, ADHD-I = 122, ADHD-HI = 12, ADHD-C = 249, Total = 929 | ReHO, ALLF and RSN | LRC (best) | 54% ADHD Subtype: 67% |
| D. Fair et al. (2013)         | ADHD                      | rsfMRI                    | TDC = 455, ADHD-I = 80, ADHD-C = 112, Total = 647 | Graph based features based on FC | SVM-based MVPA | 63.4–82.7%       |
| Deshpande, Wang, Rangaparakash, and Wilamowski (2015) | ADHD                      | rsfMRI                    | TDC = 744, ADHD = 433, Total = 1,177 | Directional connectivity measures | ANN               | 90%              |
| Semrud-Clikeman et al. (1996) | ADHD/dyslexia             | sMRI                      | HC = 10, ADHD = 10, Dyslexia = 10, Total = 30 | Morphometric measures of ROIs | Discriminant function analysis | 60.0–87%         |
| Peng, Lin, Zhang, and Wang (2013) | ADHD                      | sMRI                      | HC = 55, ADHD = 55, Total = 110 | CT measures | ELM               | 90.20%           |
| Lim et al. (2013)             | ADHD                      | sMRI                      | HC = 29, ASD = 19, ADHD = 29, Total = 77 | Voxel-wise GM volumetric measures | Gaussian process classifier | 68.2–85.2%       |
| Johnston et al. (2014)        | ADHD                      | sMRI                      | HC = 34, ADHD = 34, Total = 68 | WM maps | SVM               | 93%              |
| Igual et al. (2012)           | ADHD                      | sMRI                      | HC = 39 ADHD = 39, Total = 78 | Caudate nucleus volumetric measures | Adaboost and SVM | 72.50%           |
| Chang, Ho, and Chen (2012)    | ADHD                      | sMRI                      | HC = 226, ADHD = 210, Total = 436 | Texture features based on isotropic local binary patterns on three orthogonal planes | SVM               | 69.90%           |
| Bansal et al. (2012)          | ADHD                      | sMRI                      | HC = 42, ADHD = 41, Total = 83 | Surface morphometric measures | Semi supervised (hierarchical Clustering) | 91.00%           |
| Study                  | Diagnosis | Imaging | Data Description                                                                 | Methods | Accuracy |
|-----------------------|-----------|---------|----------------------------------------------------------------------------------|---------|----------|
| A. Anderson et al. (2014) | ADHD      | sMRI + rsfMRI + phenotypic data | TD = 472, ADHD = 276, Total = 748 Curvature index, folding index, Gaussian curvature, gray matter volume, mean curvature, surface area, thickness average, and thickness standard deviation along with functional connectivity measures and phenotypic data | NMF + decision tree | 66.80%   |
| Bohland, Saperstein, Pereira, Rapin, and Grady (2012) | ADHD      | sMRI + rsfMRI | TDC = 491, ADHD = 285, Total = 776 Various anatomical, network and nonimaging measures | SVM | 80.0% (AUC) |
| Iannaccone et al. (2015) | ADHD      | sMRI and fMRI task (flanker/NoGo) | HC = 18, ADHD = 18, Total = 36 Whole brain GLM coefficients and GM maps from VBM | SVM | 61.1-77.8% |
| Dai, Wang, Hua, and He (2012) | ADHD      | sMRI and rsfMRI | TCD = 402, ADHD = 222, Total = 624 Cortical thickness and GM maps from sMRI and ReHo and FC from rsfMRI | SVM and multi-kernel learning | 61.50% |
| Colby et al. (2012) | ADHD      | sMRI and rsfMRI | TD = 491, ADHD = 285, Total = 776 Morphological measures, FC, power spectra and graph measures | Multiple SVM | 55% |
| L. Shao, Xu, and Fu (2018) | ADHD      | rsfMRI | Dataset 1: ADHD = 22, HC = 61; Dataset 2: ADHD, 24, HC = 61; Dataset 3: ADHD = 76, HC = 118; Dataset 4: ADHD = 98, HC = 179, Total = 639 FC measures using 116 ROIs from AAL atlas | Bi-objective SVM | 81.25% (best) |
| Wolfers et al. (2016) | ADHD      | Task fMRI (stop signal task) | ADHD = 184, unaffected sibling = 103, HC = 128, Total = 415 Activation maps derived from three task regressor | GPC | 64% (AUC) |
| Cicek, Akan, and Metin (2018) | ADHD      | sMRI | ADHD = 15, HC = 11, Total = 26 Multiple structural features | KNN and naive Bayes | 100% |
| Y. Li et al. (2016) | ADHD      | rsfMRI | ADHD = 23, HC = 45, Total = 68 ROI time-series | Kernel ELM | 96.06% |
| Qureshi, Jo, and Lee (2017) | ADHD      | rsfMRI | AHDH-I = 30, ADHD-C = 30, TDC = 30, Total = 90 Global connectivity | H-ELM | 71.11% |
| Miao and Zhang (2017) | ADHD      | rsfMRI | ADHD = 82, HC = 72, Total = 154 fALFF | SVM | 98.04% |
| Yao et al. (2018) | ADHD      | rsfMRI | Dataset 1: ADHD = 112, HC = 77; Dataset 2: ADHD = 34, HC = 28, Total = 251 FC measures | Ensemble learning | 80-86% |
| Y. Zhang, Tang, Chen, Zhou, and Wang (2018) | ADHD      | rsfMRI | Dataset 1: ADHD = 25, HC = 23; Dataset 2: ADHD = 22, HC = 61; Dataset 3: ADHD = 24, HC = 62; Dataset 4: ADHD = 118, HC = 98, Total = 433 FC measures | SVM | 75% |
In general, the existing SZ predictome studies mostly used functional and structural MRI data as features and LOO-CV approach as CV, with many studies having a very small sample size. Therefore, future studies examining predictive modeling with more robust CV approaches and greater small size are necessary. Further, while these initial results suggest that SZ can be predicted with higher accuracies, the accuracy range varies substantially across these studies, and replication studies are required to confirm generalizability.

### 3.3 Major depression/bipolar disorder

While the overlapping symptoms make it challenging to differentiate between MDD and BP as well as with other disorders (e.g., SZ and schizoaffective disorder), recent studies have reported successful diagnostic prediction of BP and MDD (Arribas et al., 2010; Castafreda et al., 2011; Lueken, Hilbert, Wittchen, Reif, & Hahn, 2015; Redlich et al., 2014). We reviewed 61 studies that used neuroimaging for automatic diagnose BP and MDD which are listed in Table 2. Among the surveyed studies, structural MRI was primarily used to predict BP patients, although only few studies included BP samples. Therefore, further investigation is required prior to recommending particular classifiers or machine learning frameworks as diagnostic tools for BP prediction. As presented in Table 2, many studies have performed classification on MDD samples. While there is an increase in studies predicting MDD using rsfMRI, sMRI, and task-based fMRI as features and primarily SVM classifier, with the accuracies ranging from 52% to 99%, the sample size is relatively small in most studies (Table 2).

1 **Structural MRI**: Castafreda et al. (2009) utilized sMRI data where they used gray matter maps as features from 37 MDD and 37 HC participants and achieved 67.6% classification accuracy. Another recent study with relatively large sample size (i.e., 62 MDD and 62 HC) used voxel-based gray matter maps driven from sMRI and SVM classifier and achieved 90% accuracy (Mwangi et al., 2012). Few recent studies have also focused on cross-disorder prediction of MDD and BP, as well as MDD and SZ. For instance, using multivariate patterns form gray matter differences and LOO-CV, Redlich and colleagues differentiated MDD from BP patients with 79.3% accuracy, and further validated the findings using an independent dataset and test-retest validation with 65.5% accuracy (Redlich et al., 2014).

2 **Functional MRI**:  
   
   a **Task-based**: Among the initial MDD prediction studies are two task-based fMRI studies by Marquand et al. (2008) (verbal memory N-back task, 67.5% accuracy) and Fu et al. (2008) (sad facial processing task, 84% accuracy). Other task-based MRI studies include the works performed by Sato et al. (2015) (task paradigm included written statements of actions counter to social and moral values described by social, 78.3% accuracy), Johnston, Tolomeo, et al. (2015) (instrumental loss-avoidance and win–gain reinforcement learning...
| References                  | Disorder | Modality | Number of subjects | Feature type                                      | Classifier | Overall accuracy |
|-----------------------------|----------|----------|--------------------|--------------------------------------------------|------------|------------------|
| Soriano-Mas et al. (2007)   | OCD      | sMRI     | OCD1 = 72 HC2 = 72 | Mean difference value of OCD and controls        | N.S.       | 76.6             |
| Weygandt et al. (2012)      | OCD      | Task fMRI| OCD = 10 HC = 10,  | Voxel-based feature set and                       | SVM        | 100              |
| Shenas, Halici, and Cicek (2014) | OCD     | rsfMRI; task fMRI | OCD = 12 HC = 12, Total = 24 | Region-based feature set and                       | LDC; SVM   | 74               |
| Li et al. (2014)            | OCD      | dMRI     | OCD = 28 HC = 28,  | Voxel-based feature set                           | SVM        | 84               |
| Shenas, Halici, and Cicek (2013) | OCD     | rsfMRI   | OCD = 12, HC = 12, Total = 24 | FC measures                                       | SVM        | 66%              |
| Gruner et al. (2014)        | OCD      | rsfMRI   | OCD = 23, HC = 23,  | ICA components                                    | LR         | 80.10%           |
| Parrado-Hernández et al. (2014) | OCD | sMRI   | OCD = 86, HC = 86, Total = 172 | Voxel-based measures                               | SVM        | 73.87%           |
| Yun, Jang, Kim, Jung, and Kwon (2015) | OCD | sMRI | OCD = 56, HC = 75, Total = 131 | Structural covariance                             | SVM        | 90.7–95.6%       |
| X. Hu et al. (2016)         | OCD      | sMRI     | OCD = 33, HC = 33,  | Voxel-based measures                               | SVM        | 81.82%           |
| Sen et al. (2016)           | OCD      | rsfMRI   | OCD = 16, HC = 13,  | Network measures                                   | SVM        | 80%              |
| Takagi et al. (2017)        | OCD      | sMRI     | Main dataset OCD = 56, HC = 52; validation: OCD = 10, HC = 18, Total = 136 | FC measures | Sparse LR | 73%              |
| Trambaioli, Biazoli Jr, Balardin, Hoexter, and Sato (2017) | OCD      | rsfMRI   | OCD = 38, HC = 36, Total = 74 | Regional measures                                  | SVM        | 71.64%           |
task, 97% accuracy [best result]), and Shimizu et al. (2015) (semantic and phonological verbal fluency task, 95% accuracy [best result]).

b Resting-state: Among the rsfMRI studies, Craddock, Holtzheimer III, Hu, and Mayberg (2009) used region-based feature set from rsfMRI and achieved 95% accuracy. Further, using rsfMRI data and SVM classifier, Yu, Shen, Zeng, et al. (2013) achieved 84.2% accuracy.

c Diffusion MRI: Studies utilizing features from dMRI data include SVM classifier based pipelines and features such as anatomical connectivity, graph metric of white matter connectivity, and FA, with classification accuracies ranging from 63% to 91% (Deng et al., 2018; Fang et al., 2012; Sacchet, Prasad, et al., 2015; Schnyer et al., 2017).

d Multimodal: Using sMRI and dMRI data and LDC classifier, Ota et al. (2013) obtained about 80% accuracy, to distinguish MDD from SZ and HC individuals. Another large-scale MDD study (total sample size = 307) by J. Yang et al. (2018) used structural and dMRI data and reported 75% accuracy.

While these studies utilized a variety of MRI based features, such as sMRI, resting and task-based fMRI and dMRI, they provide evidence of brain-based differential diagnoses of MDD and BP. Future studies should employ large-scale sample size, and more BP prediction research.

3.4 | Autism spectrum disorder

ASD is a neurodevelopmental disorder characterized by impaired social communication, deficits in social-emotional reciprocity, deficits in nonverbal communicative behaviors used for social interaction and stereotypic behavior. Since 2010, a few studies have investigated automatic diagnosis of ASD in both male-only and male–female samples (Ecker, Marquand, et al., 2010; Ecker, Rocha-Rego, et al., 2010; Jiao et al., 2010) with promising results showing accuracies ranging from 81% to 90% (Table 3). We surveyed 30 papers in automatic diagnosis of ASD using MRI-based features which are listed in Table 3.

1 Structural MRI: Using voxel-based feature set (Ecker, Rocha-Rego, et al., 2010) and region-based feature set (Ecker, Marquand, et al., 2010) from sMRI data and SVM classifiers, Ecker and colleagues predicted ASD with 81 and 90% accuracies, respectively. Interestingly, Uddin and colleagues employed a searchlight algorithm to sMRI features, where a small number of voxels within the spatial proximity with one another provide the predicting features (Uddin et al., 2011). The resulting model obtained 92% classification accuracy for ASD diagnosis.

2 Functional MRI:
(a) Task-based: More recently, features from task-based fMRI and SVM, LRC, LDC, and GPC classifiers were used for ASD prediction, with accuracies ranging from 70% to 96% (Table 3).
(b) Resting-state: Other ASD prediction studies include region-based feature set and automatic feature selection from resting fMRI data with 79% accuracy (J. S. Anderson et al., 2011) and ICA component based features from resting fMRI data with 78% accuracy (Uddin et al., 2013).

3 Diffusion MRI: Ingalhalikar and colleagues used region-based feature set and automatic feature selection from dMRI data and achieved 80% accuracy (Ingalhalikar et al., 2011). Another dMRI-based prediction study by Zhang and colleagues used SVM classifier on FA and mean diffusivity measures and reported to achieve 78% accuracy.

4 Multimodal: Studies utilizing multimodal MRI data for ASD prediction include the works by Deshpande and colleagues where both task fMRI and dMRI data were used to identify causal connectivity weights, connectivity values, and FA features, resulting in 95.9% accuracy (Deshpande et al., 2013). Another work discriminating between ASD and HC participants used rsfMRI and sMRI based features and achieved 65% accuracy (Ghiassian et al., 2016).

While these studies have utilized all available MRI data modalities as features for predicting ASD, the number of sample size as well as studies based on each modality are relatively low. Additionally, heterogeneity associated with ASD, including subtypes of lower versus higher functioning patients, should be further investigated using predictive models.

### TABLE 6 | Social anxiety disorder

| References                  | Disorder | Modality | Number of subjects | Feature type          | Classifier | Overall accuracy |
|-----------------------------|----------|----------|--------------------|-----------------------|------------|-----------------|
| F. Liu, Guo, et al. (2015)  | SAD      | rsfMRI   | SAD = 20, HC = 20, Total = 40 | Region-based feature set | SVM        | 82.5%           |
| Frick et al. (2014)         | SAD      | Task fMRI; sMRI | SAD = 14, HC = 12, Total = 26 | Voxel-based feature set | SVM        | 84.5%           |

### TABLE 7 | Posttraumatic stress disorder

| Reference                  | Disorder | Modality | Number of subjects | Feature type          | Classifier | Overall accuracy |
|-----------------            |----------|----------|--------------------|-----------------------|------------|-----------------|
| Q Gong et al. PTSD (2014) | PTSD     | sMRI     | PTSD earthquake = 50, no-PTSD earthquake = 50, HC = 40, Total = 140 | Voxel-based feature set | SVM        | No-PTSD vs. PTSD: 76.00, No-PTSD vs. HC: 85.00, PTSD vs. HC: 91.00 |
3.5 | Attention-deficit/hyperactivity disorder

One of the most commonly found neurodevelopmental disorders is ADHD. However, given the lack of biological-based diagnosis approach, ADHD is currently diagnosed based on behavioral symptoms only. In this review, we surveyed 35 papers in automatic diagnosis of ADHD using MRI-based features which are listed in Table 4.

1 Structural MRI: Studies based on sMRI features reported accuracies for ADHD classification ranging from about 72% to 93% (Igual et al., 2012; Johnston et al., 2014; Lim et al., 2013). Using voxel-based feature set from sMRI data and automatic feature selection approach, Johnston and colleagues trained the features using a SVM classifier and obtained 93% accuracy (Johnston et al., 2014). Lim and colleagues used whole brain gray and white matter from sMRI data and GPC classifier and predicted ADHD with 79.3% accuracy (Lim et al., 2013).

2 Functional MRI:

a Task-based: In order to predict ADHD, Hart and colleagues used stop signal task-based fMRI data (Hart, Chantiluke, et al., 2014; 77% accuracy) and temporal discounting task-based fMRI data (Hart, Marquand, et al., 2014; 75% accuracy). However, these studies were performed using relatively small sample sizes, and therefore may lack generalizability across other independent samples.

b Resting-state: A recent study by Zhu and colleague used voxel-based feature set and automatic feature from rsfMRI data and LDC and SVM classifiers to classify ADHD with an accuracy of 75–80% (C.-Z. Zhu et al., 2008). Other resting-state studies include the work by Sidhu and colleagues where they used SVM classifier in a large sample with 668 participants and obtained 76% accuracy (best result; Sidhu et al., 2012).

c Diffusion MRI: While studies have reported disrupted white matter tracts in ADHD patients (van Ewijk, Heslenfeld, Zwiers, Buitelaar, & Oosterlaan, 2012), to date no published studies have examined dMRI based features for ADHD prediction.

d Multimodal: Several multimodal studies for ADHD prediction include sMRI, rsfMRI, and phenotypic data (A. Anderson et al., 2014), sMRI and rsfMRI data (Bohland et al., 2012; Colby et al., 2012; Dai et al., 2012), and sMRI and task-based MRI data (Iannaccone et al., 2015), with accuracy ranging from 55% to 80%.

3.6 | Obsessive–compulsive disorder

Only a few recent studies have applied classification algorithms to OCD. We surveyed 12 papers focused on automatic diagnosis of OCD using MRI-based features which are listed in Table 5.

1 Structural MRI: Using an approach to calculate the distance between individual participants and the mean of OCD and control group based on sMRI derived measures, Soriano-Mas and colleagues...
distinguished between OCD patients and HCs (Soriano-Mas et al., 2007). Other sMRI studies for OCD prediction reported overall accuracies ranging from 73% to 95.6% (X. Hu et al., 2016; Parrado-Hernández et al., 2014; Takagi et al., 2017; Yun et al., 2015).

2 Functional MRI:

a Task-based: In a task-based fMRI study presenting emotional valence stimuli, OCD patients were classified with perfect accuracy in a very small sample of 10 participants per group (Weygandt et al., 2012).

b Resting-state: The surveyed rsfMRI studies mostly used SVM classifier to predict OCD (Sen et al., 2016; Shenas et al., 2013; Trambaioli et al., 2017; accuracy ranging from 66% to 80%), with an exception of logistic regression classifier (Gruner et al., 2014; 80% accuracy).

c Diffusion MRI: Li and colleagues used voxel-based feature set from dMRI data and SVM classifier to classify OCD with 84% accuracy (Li, Tran, et al., 2014).

d Multimodal: Using features from multimodal resting-state MRI and task MRI data Shenas and colleagues reported 74% accuracy for OCD prediction while applying SVM and LDC classifiers (Shenas et al., 2014).

3.7 Social anxiety

To date, only two studies on SAD have been published which included relatively small samples, with accuracies above 80% (Frick et al., 2014; Liu et al., 2015). These studies derived multivariate patterns from different MRI modalities suggesting that features relevant for SAD
classification can be extracted and analyzed across modalities. Both studies reported useful features to be distributed across widespread brain areas, rather than localized brain regions typically associated with anxiety. We surveyed two papers focused on automatic diagnosis of SAD using MRI-based features which are listed in Table 6.

3.8 | Posttraumatic stress disorder

Only one study to date has performed discriminative analysis on PTSD, where 50 earthquake survivors with and without PTSD were compared to controls using structural imaging (Q Gong et al., 2014). Patients with PTSD were classified with an accuracy of 91%, with the most discriminative features found in different brain areas, particularly in left and right parietal regions. Table 7 presents the surveyed paper in automatic diagnosis of PTSD using MRI-based features.

3.9 | Substance dependence

To date, only a few predictive studies on SD (e.g., alcohol, nicotine, and cocaine addiction) and treatment completion have been performed, with only one study implementing a multimodal imaging approach to predict alcohol consumption and treatment effects in the
brain (Cosa et al., 2017), although using data from Marchigian-Sardinian (msP) rats. Table 8 provides the seven surveyed paper in automatic diagnosis of SD using MRI-based features.

1 Structural MRI: Using SVM-RFE classifier, Ding and colleagues utilized mean gray matter volume features from sMRI data to predict nicotine dependence with 69.6% accuracy (Ding et al., 2015). Alcohol dependence was predicted in a recent study using regional gray matter maps from sMRI and weighted robust distance and SVM classifiers (Guggenmos et al., 2018), resulting in 74% classification accuracy.

2 Functional MRI:
   (a) Task-based: Only one study investigated heroin-dependence using FNC measures from task-based fMRI and SVM classifiers, and achieved 80.58% accuracy (Steele et al., 2018).
   (b) Resting-state: Using FC measures as features from rsfMRI data and SVM classifiers, Pariyadath and colleagues predicted nicotine dependence with a maximum accuracy of 78.6% (Pariyadath et al., 2014). Ding and colleagues further used rsfMRI data and SVM based classifiers to successfully predict nicotine dependence (Ding et al., 2017; local measures and network measures, 75.5% accuracy). Alcohol Dependence was predicted in a recent study using rsfMRI based FC measures and Random Forest classifier, obtaining 87% accuracy (Zhu, Du, et al., 2018). Using connectivity measures from rsfMRI data, SVM classifier was used to predict nicotine dependence with 95.2% accuracy (Vergara et al., 2017).

3.10 | Analysis of the survey

Figure 3 demonstrates some key features from this survey. Figure 3a presents the number of studies published in each year for each modality, Figure 3b shows the number of studies published in each year for
each disorder type, and Figure 3c shows the number of studies published in each year for each disorder type and each modality. These figures show that there is a publication growth since 2007, and since 2010, the number of studies has been growing rapidly. Only one ADHD prediction study was performed before 2000, which included a sample size relatively low compared to the recent ADHD publications (Semrud-Clikeman et al., 1996). Particularly, all major disorders’ studies (i.e., SZ, MDD/BP, ASD, and ADHD) have shown a peak between 2011 and 2013, which might be due to the recent data-sharing initiatives. From Figure 3c, it can be observed that all MRI-based features have been utilized for prediction of these major disorders, with a few multimodal studies. Further, it shows that structural MRI (sMRI) is the most popular choice of modality, particularly for SZ studies. For MDD/BP, ASD, and ADHD studies, rsfMRI is the most popular modality. Moreover, compared to dMRI, multimodal studies are more common across these major disorders.

Figure 3d shows the overall prediction accuracy against the commonly used classifiers in each disorder type, Figure 3e reports the overall prediction accuracy against each modality and each disorder type, and Figure 3f presents the total sample size against each disorder type for each modality. SVM classifier was the most popular across all major disorders, followed by LDA.

Figure 4a shows the overall accuracy against the total sample size for each disorder and for each classifier used in the studies, Figure 4b shows the overall accuracy against the total sample size for each modality and for each classifier used in the studies, and Figure 4c shows the overall accuracy against the total sample size for each disorder and each modality used in the studies. Interestingly, even with a sample size smaller than 100, almost all studies reported very high accuracies, with few reporting...
100% classification accuracy. The reported overall accuracy decreases with increasing sample size in most of the major disorders, including SZ and MDD/BP, which is concerning as it indicates that the classification framework utilized in these small sample size studies might not be generalized across large-scale studies. Further, SVM classifier shows to achieve very high accuracy across most of the major disorders. Also, rsfMRI and task-based SZ, MDD/BP and ADHD studies show high accuracy rate.

Figure 4d reports the sample size distribution for number of studies in this survey. The dashed lines represent mean (red) sample size across all studies. Figure 4e shows the overall accuracy against each modality for each disorder and for each classifier used in the studies. MDD/BP and ADHD studies show high accuracy rate.

Based on our surveyed studies (Tables 1–8), volume and cortical thickness from structural MRI, the activation maps and functional connectivity among ROIs or ICA components from fMRI data and FA from dMRI data were some of the most commonly used features for classification. Further, SVM was the most popular classifier of choice across all disorders’ studies, and different extensions of SVM methods, including linear, nonlinear (different kernel), and SVM with recursive feature elimination (SVM-RFE), were observed. Other popular classifiers from the surveyed studies include LDA, Gaussian process classifier (GPC) and random forest (RF).

Figures 5 and 6 present the cumulative density function (CDF) for (a) publication year per modality, (b) sample size per modality, (c) accuracy per modality, and (d) accuracy per classifier, for the main disorders (disorders with very few number of studies were excluded). An interesting observation from this summary is that, there is a growth in multimodal prediction studies for many of these major mental disorders.

4 | TRANSLATIONAL PERSPECTIVE OF BRAIN-BASED PREDICTOME RESEARCH FOR CLINICAL APPLICATION

4.1 | Translating predictive outcomes toward clinical utility

Typically, in a research-based setup, predictive studies are implemented using two or more well-proportioned groups of patients with mental illness and their healthy counterparts. The group labels are carefully diagnosed before training a supervised classification algorithm, where exclusion of subjects with uncertain diagnoses or comorbidities is a common practice (Wolfers et al., 2015). However, in case of real clinical population, the disease diagnosis is a more complex and sophisticated process. Thus, considerable improvement in the field of predictive modeling is required before they will be useful to apply in clinical practice. In many clinical cases, the central question to be addressed is not as simple as how to distinguish patients from controls, but rather the specific distinction between different illnesses in the same population (i.e., subtypes). Simply put, a differential diagnostic process is required before accurate clinical implementation of the available tools can be made available. Moreover, another limitation of the current predictive modeling approaches is the lack of appropriate (or any) identification of comorbidities among patients, which is essential to properly assign memberships across multiple diagnostic classes. To date, there are only a few existing studies in the current literature across different disorders have demonstrated multi-class classification (i.e., subtype classification) of three or more disorders (Costafreda et al., 2011; Du et al., 2015; Dwyer et al., 2018; Grotegerd et al., 2013; Koutsouleris et al., 2015;
Lim et al., 2013; Redlich et al., 2014; Serpa et al., 2014; Yu, Shen, Zeng, et al., 2013). However, to the best of our knowledge, no such studies have adequately addressed the issue of comorbidity. Although existing limitations should be addressed before clinical application of the research-based diagnostic/predictive tools become available, even current approaches can be used as a supplement to clinical diagnosis with higher decision-making uncertainty or even to make decisions for any follow-up medical procedure requirements. Beyond the traditional subjective approaches, incorporating quantitative prediction approaches of psychiatric disorders can assist more accurate clinical diagnosis (Krueger et al., 2018). However, there are several key factors that need to be considered prior to a successful clinical translations of the neuroimaging predictive modeling, including reliability of neuroimaging-based findings and subject-level clinical transition (Saggar & Uddin, 2019). Moreover, some of the existing gaps in psychiatric neuroimaging research need to be addressed (Etkin, 2018, 2019).

More specifically, some immediate clinical application of predictive modeling may include:

- **Diagnosis based on "biomarkers" in addition to clinical interviews:** Current clinical diagnoses are primarily made based on clinical interviews. In addition to the existing behavioral measures,
predictive modeling outcomes can provide more accurate brain-based endophenotypes or "biomarkers", and combining both measures can provide deeper understanding of biological pathways, mechanisms and progression of the disease. For example, brain-based biomarkers for ASD and ADHD could aid in disease diagnosis more objectively (Uddin, Dajani, Voorhies, Bednarz, & Kana, 2017).

- **Multi-class classification**: In clinical practice, it is very challenging to diagnose severe mental disorders that share substantial overlapping symptoms such as SZ, schizoaffective, unipolar and bipolar depression, and mood disorders. By obtaining estimates of predictive confidence level across multiple disease classes, predictive models could offer to identify comorbidities in patient groups. Recent multi-class approaches can specify a predictive value for all classes of interest for each subject, and are able to indicate comorbidity of subjects for multiple disorders based on their predictive confidence level (Filippone et al., 2012). Other approaches to identify comorbidities include multi-label classification (M.-L. Zhang & Zhou, 2014) and multi-task learning (Pan & Yang, 2010), where multiple class labels are assigned to each sample.

- **Patient screening**: Before proceeding with clinical procedures or expert opinions, neuroimaging-based predictive outcomes can be
utilized as a patient screening phase. A prior screening of patients using predictive modeling might reduce the time and expenses related to clinical interviews.

- **Treatment response/outcome prediction**: In addition to aiding the decision-making procedure in clinical diagnoses, predictive techniques can also be utilized in predicting treatment response (Bzdok & Meyer-Lindenberg, 2018) (e.g., Gong et al., 2011) and treatment outcome (e.g., Schmaal et al., 2015). By monitoring treatment outcomes and pursuing potential treatments using predictive modeling, the clinical diagnosis can become more cost-efficient.

- **Drug trial design**: Based on predictive modeling outcomes, future response can also be classified. By selecting subsets of individuals who are most likely to respond to a particular medication, more efficient drug trials can be designed. For example, medication-class of response to mood stabilizers (bipolar) or antidepressants (depression) can be classified using machine learning approaches (Osuch et al., 2018).

### 4.2 Prediction of continuous measures versus categorical diagnoses

Most of the mental illness prediction studies surveyed in this review are based on assignment of discrete or categorical class labels for test samples. However, the categorical diagnosis approach overlooks the continuous measures while predicting a certain disease class, which can lead to misleading outcomes, or miss sub-clinical tendencies that can be useful for predicting risk. For more reliable outcomes, predictive modeling using continuous measure, such as pattern regression, can become a valuable tool. Moreover, for mental illness prediction using brain-based features, regression-based modeling can be used to estimate the disease progression and treatment outcomes, and can estimate continuous measures (e.g., neuropsychiatric or cognitive measures). In order to estimate continuous clinical measures from neuroimaging data, a recent study by Wang and colleagues proposed a framework using the relevance vector machine (RVM) to build regression and obtained higher classification accuracy and...
better generalizing ability compared to support vector regression (Y. Wang, Fan, Bhatt, & Davatzikos, 2010). Another study explored inter-regional cortical thickness correlations to identify and characterize the autism diagnostic observation schedule score in ASD (Sato et al., 2013). The results from this study showed that structural covariance measures among multiple brain networks are associated with autistic symptoms. Further, Tognin and colleagues used relevance vector regression to Predict Positive and Negative Syndrome Scale scores of subjects at high risk of psychosis based on gray matter volume and cortical thickness measurements (Tognin et al., 2014). More recently, studies have started to predict continuous measures of assessment in both health and disease (e.g., the research domain criterion [RDoC]; T. Insel et al., 2010). These studies suggest that promising results can be achieved by using continuous measure for disease prediction in addition to categorical diagnosis.

4.3 Prediction of disease risk (prodromal state)

While challenging, early diagnosis of individuals at high risk of future mental illness is very critical in order to delay or prevent the disease progression. Since most mental illnesses typically have an onset in adolescence or early adulthood (Kessler et al., 2005), early detection could delay, or even prevent, future onset of these severe illnesses in high-risk adolescents. Predictive modeling based approaches offer promising tools to be used for clinical diagnosis, such as identification of neuroimaging-based biomarker that can support early identification of potentially at-risk individuals of developing mental disorders, with the potential risk being unidentified. However, only a limited number of studies have explored the discriminative power of predictive modeling to identify healthy adolescents genetically at-risk for mental illnesses. Mourão-Miranda and colleagues investigated whether Gaussian process classifiers (GPC) based machine learning algorithm using neuro-imaging data could help differentiate healthy adolescents genetically at-risk for BP disorder and other Axis I psychiatric disorders from healthy adolescents at low risk of developing these disorders, and identify those healthy genetically at-risk adolescents who were most likely to develop future Axis I disorders (Mourão-Miranda et al., 2012). Results from this study showed that the model can discriminate healthy low-risk from healthy adolescents genetically at-risk for Axis I disorders, as well as help predict which at-risk adolescents subsequently develop

**FIGURE 11** A joint estimation approach for schizophrenia prediction using parallel group ICA + ICA. Flowchart showing steps to extract first-level fMRI (a) and sMRI (b) features, feature integration using parallel GICA+ICA (c), group-level components extraction from GICA (d), subject-level GICA components extraction (e), group-level GICA components extraction (f), FNC analysis. Figures reprinted with permission from Qi et al. (2019)
these disorders. Further, Guo and colleagues used fractional amplitude of low-frequency fluctuation (fALFF) to characterize signals from the default-mode network in unaffected siblings of SZ patients using resting-state functional MRI data (W. Guo, Su, et al., 2014). Also, using machine learning methods, Fan and colleagues explored structural endophenotypes in unaffected family members of SZ patients, and reported that family members had structural profiles highly overlapping with those of SZ patients (Fan et al., 2008).

4.4 | Prediction of disease onset and treatment outcome/responses

Typically, prediction approach is applied for predicting disease onset, progression, diagnosis, or treatment outcome/response. For more clinical context, predictive modeling can be used as a diagnostic tool or for predicting treatment response. In order to predict the onset of MDD, a recent machine learning-based longitudinal, 5-year follow-up study was performed (Foland-Ross et al., 2015), where cortical thickness measures were used to predict differences between participants who later experienced depression and subjects who remained healthy, with a predictive accuracy of 70% ($p = .021$). A few studies have attempted to predict the treatment response of mental disorders. Using machine learning approach, Liu and colleagues obtained an accuracy of 82.9% to predict treatment response and resistance (F. Liu et al., 2012). Another study by Osuch and colleagues examined the translational perspective of fMRI (i.e., spatially independent components [ICs]) based classification for medication-class of response prediction in patients with MDD and BP (Osuch et al., 2018). Using SVM classifiers and a nested CV approach, the algorithm predicted the medication class of response 91.7% of the time. To prevent or delay disease progression, as well as to develop optimized treatment plans, further applications of machine learning-based disease onset and treatment response prediction are required.

5 | CURRENT TRENDS IN THE BRAIN-BASED PREDICTOME

5.1 | Univariate versus multivariate approaches

The recent methodological developments of a wide range of multivariate neuroimaging analysis approaches have received growing attention due to their ability to examine additional features beyond the voxel-wise, univariate techniques. In case of univariate, functional neuroimaging approaches, typically a model of the expected response variable is fitted independently at each voxel's time course, and then further tests are performed using the estimated response levels during experimental conditions (K. J. Friston et al., 1994). Although convenient, this approach introduces a large number of statistical tests as it results in statistical difference maps of responses by targeting only selective brain locations related to a specific stimulus (i.e., voxels or ROIs; K. J. Friston et al., 1994). Thus, it limits the study of stimulus-
independent relationships among brain regions (e.g., functional connectivity). In addition, the univariate approach does not allow estimation of the stimulus effects at multiple brain locations. Multivariate neuroimaging approaches take into account the full spatial pattern of brain activity simultaneously at multiple spatial locations, and are able to detect subtle but localized measures of brain activity that are not captured by univariate approaches (Allen et al., 2011; Habeck et al., 2008; Habeck et al., 2005; J. Liu & Calhoun, 2014; McIntosh & Lobaugh, 2004; Moeller & Strother, 1991; Narayanan et al., 2015; Sui, Adali, Yu, Chen, & Calhoun, 2012). In contrast to the traditional univariate, model-based approach which lacks the ability to directly address interaction between voxels/regions, multivariate approach estimates correlation or covariance of activation across brain regions. Multivariate results can also be more reliably translated as a signature of underlying brain networks. Recent multivariate neuroimaging methods offer to analyze the relationship between a stimulus and the responses simultaneously measured at many locations, such as spatial response patterns or multi-voxel response patterns. Compared to univariate techniques, multivariate approaches provide potentially greater statistical power and better reproducibility.

In neuroimaging-based diagnoses, multivariate machine learning methods integrate available features simultaneously to jointly differentiate between groups. Typically, for a multivariate machine learning approach, a classifier is trained on a training dataset to predict different classes (e.g., patient groups), which is then applied to the testing dataset. The classification accuracy (i.e., correct classification rate) is then estimated, using a CV procedure for improved accuracy. To date, a number of multivariate machine learning methods have been applied in neuroimaging-based prediction, such as SVM, k-nearest neighbor (KNN), Gaussian Naive Bayes (GNB), and LDA. Among these classifiers, the SVM-based classifier has been reported to obtain greater classification performance (Davatzikos et al., 2005; Mitchell

**FIGURE 13** An approach to classify schizophrenia (SZ) using resting-state functional network connectivity (FNC) measures. (a) Feature extraction and classification steps of the FNC-based framework. (b) group-wise mean FNC (left), and T values (FDR corrected, \( p < .05 \)) (right). (c) Group difference in mean FNC measures (left), and FDR corrected (\( p < .05 \)) T values showing group difference. Figures reprinted with permission from Arbabshirani et al. (2013)
| References                        | Disorder                      | Disorder Type | Number of subjects | Feature Type                                                                 | Classifier          | Overall Accuracy |
|----------------------------------|-------------------------------|---------------|--------------------|-----------------------------------------------------------------------------|----------------------|------------------|
| Calhoun et al. (2008)            | SZ/BP                         |               | HC = 26, SZ = 21, BP = 14, Total = 61 | Network maps (DMN and temporal lobe) extracted by ICA                      | ED                   | 83–95%           |
| Demirci, Clark, and Calhoun (2008) | SZ                            |               | HC = 36, SZ = 34, Total = 70 | Network maps extracted by ICA                                               | Projection pursuit   | 80–90%           |
| Yang et al. (2010)               | SZ                            |               | HC = 20, SZ = 20, Total = 40 | Multiple features include SNP, voxels in fMRI map, and components of fMRI activation | SVM                  | 74–87%           |
| Castro et al. (2011)             | SZ                            |               | HC = 54, SZ = 52, Total = 106 | Activation maps extracted by GLM and networks extracted by ICA             | SVM                  | 95%              |
| Fan et al. (2011)                | SZ                            |               | HC = 31, SZ = 31, Total = 62 | Network maps extracted by ICA                                               | SVM                  | 87%              |
| Bassett et al. (2012)            | SZ                            |               | HC = 29, SZ = 29, Total = 58 | FC between 90 ROIs (AAL atlas)                                             | SVM                  | 75%              |
| W. Du et al. (2012)              | SZ                            |               | HC = 28, SZ = 28, Total = 56 | Network maps extracted by ICA                                               | Fisher discriminant function | 93–98%           |
| Tang et al. (2012)               | SZ                            |               | HC = 22, SZ = 22, Total = 44 | FC between 90 ROIs (AAL atlas)                                             | SVM                  | 93.20%           |
| Venkataaram et al. (2012)        | SZ                            |               | HC = 18, SZ = 18, Total = 36 | FC between 77 anatomical ROIs                                               | RF                   | 75%              |
| A. Anderson and Cohen (2013)     | SZ                            |               | HC = 29, SZ = 19, Total = 48 | Graph measures                                                             | SVM                  | 80%              |
| Anticevic et al. (2013)          | SZ/BP                         |               | HC = 90, SZ = 90, BP = 47, Total = 227 | Thalamus seed-based connectivity                                           | SVM                  | 73.90%           |
| Arbabshirani et al. (2013)       | SZ                            |               | HC = 28, SZ = 28, Total = 56 | FNC between independent components extracted by ICA                       | Different linear and nonlinear methods | 63–96%           |
| Su et al. (2013)                 | SZ                            |               | HC = 32, SZ = 32, Total = 64 | FC between 116 ROIs (AAL atlas)                                            | SVM                  | 82.80%           |
| Yu, Shen, Zeng, et al. (2013)    | SZ                            |               | HC = 38, SZ = 32, Total = 70 | FC between 116 ROIs (AAL atlas)                                            | SVM                  | 80.90%           |
| Yu, Shen, Zhang, et al. (2013)   | SZ                            |               | HC = 25, HC sibling = 25, SZ = 24, Total = 74 | FC between 116 ROIs (AAL atlas)                                            | SVM                  | 62.00%           |
| Guo, Kendrick, Yu, Wang, and Feng (2014) | SZ                         |               | HC = 62, SZ = 69, Total = 131 | FC between 90 ROIs (AAL atlas)                                             | SVM                  | 79–82%           |
| Shen et al. (2014)               | SZ                            |               | HC = 25, SZ = 24, Total = 49 | Dynamic ALFF of ROIs                                                       | SVM                  | 81.30%           |
| Watanabe et al. (2014)           | SZ                            |               | HC = 67, SZ = 54, Total = 121 | FC between 347 spherical nodes                                             | Lasso, elastic-net, graph-net and fused lasso | 59.7–90.8%     |
| H. Cheng, Newman, et al. (2015)  | SZ/BP                         | Schizoaffective disorder | HC = 29, SZ = 19, Total = 48 | Graph measures                                                             | SVM                  | 80%              |
| Du et al. (2015)                 | SZ/BP                         | Schizoaffective disorder | HC = 24, SZ = 24, BP = 24, SZA (manic) = 24, SZA (depressive) = 13, Total = 109 | Network maps from ICA                                                  | SVM                  | 68.75%           |
| Kaufmann et al. (2015)           | SZ                            |               | HC = 196, SZ = 71, Total = 267 | FNC between independent components extracted by ICA                       | LDA                  | 84.40%           |
| Cabral et al. (2016)             | SZ                            |               | HC = 74, SZ = 71, Total = 145 | FC between 90 ROIs (AAL atlas)                                             | SVM                  | 70.50%           |
| Study                                      | Group       | HC, SZ, Total | Brain Region(s)                                      | Classification Method | Acc. (%) |
|-------------------------------------------|-------------|---------------|-----------------------------------------------------|-----------------------|----------|
| J. Kim et al. (2016)                      | SZ          | 50, 50, 100   | FC between 116 ROIs (AAL atlas)                     | DNN                   | 86%      |
| P. Mikolas et al. (2016)                  | SZ          | 63, 63, 126   | Seed-based FC                                       | SVM                   | 73%      |
| Rashid et al. (2016)                      | SZ/BP       | 61, 60, 38, 159 | Static FNC, dynamic FNC and combined static and dynamic FNC | SVM                   | 59.12–88.68% |
| Cetin et al. (2016)                       | SZ          | 44, 47, 91    | Static FNC and dynamic FNC from fMRI and MEG data   | SVM                   | 51.65–90.11% |
| Cetin et al. (2016)                       | SZ          | 348, 182, 530 | FNC between independent components extracted by ICA  | Regularized LDA       | 69–78.3% |
| Y. Liu et al. (2017)                      | SZ          | 31, 48, 79    | Voxel-mirrored homotopic connectivity                | SVM                   | 94.93%   |
| J. S. Anderson et al. (2011)              | ASD         | 40, 40, 80    | Whole brain FC between 7,266 ROIs                   |Statistic based classification score | 79% |
| Murdaugh et al. (2012)                    | ASD         | 14, 13, 27    | FC between DMN ROIs                                 | LR                    | 96.30%   |
| H. Wang, Chen, and Fushing (2012)         | ASD         | 29, 29, 58    | FC between 106 ROIs (AAL atlas)                     | LR                    | 82.80%   |
| Deshpande et al. (2013)                   | ASD         | 15, 15, 30    | FC from fMRI and FA from DTI                         | SVM                   | 95.90%   |
| Nielsen et al. (2013)                     | ASD         | 517, 447, 964 | Whole brain FC between 7,266 ROIs                   |Statistic based classification score | 60% |
| Uddin et al. (2013)                       | ASD         | 20, 20, 40    | Independent components extracted by ICA             | SVM                   | 78–83%   |
| Zhou et al. (2014)                        | ASD         | 153, 127, 280 | Graph measures                                     | SVM and Bayesian network | 70%   |
| C. P. Chen et al. (2015)                  | ASD         | 126, 126, 252 | FC between 220 ROIs from meta-analysis              | RF                    | 90.80%   |
| Iidaka (2015)                             | ASD         | 328, 312, 640 | FC between 90 ROIs (AAL atlas)                      | Probabilistic neural network | 90%   |
| Plitt et al. (2015)                       | ASD         | 59, 59, 118   | Whole brain FC from different atlas                 | SVM                   | 76.60%   |
| H. Chen et al. (2016)                     | ASD         | 128, 112, 240 | Frequency distribution-based FC                      | SVM                   | 79.20%   |
| Abraham et al. (2017)                     | ASD         | 468, 403, 871 | Whole brain FC from different atlas                 | SVM                   | 68%      |
| Jahedi, Nasamran, Faires, Fan, and Müller (2017) | ASD | 126, 126, 252 | FC between 220 ROIs from the meta-analysis of functional imaging studies | Conditional RF | 71% |
| Ktena et al. (2018)                       | ASD         | 468, 403, 871 | Graph measures                                     | CNN                   | 80%      |
| Sadeghi et al. (2017)                     | ASD         | 31, 28, 59    | Local and global functional network properties      | SVM                   | 92%      |
| Bernas et al. (2018)                      | ASD         | 18, 12, 12, 12, 24 | Time of in-phase coherence between independent component extracted from ICA | LDA and SVM | 86.70% |
|                                             | ASD         | 530, 505, 1,035 | Auto-encoder                                       |                       | 70%      |
| Study                        | Group                  | Control 1 | Control 2 | Control 3 | Control 4 | Total     | Methodology                                                                 |
|------------------------------|------------------------|-----------|-----------|-----------|-----------|-----------|-----------------------------------------------------------------------------|
| C.-Z. Zhu et al. (2008)      | ADHD                   | HC = 12   | ADHD = 12 | Total = 24 |           |           | ReHo, PCA-FDA, 85%                                                          |
| Bohland et al. (2012)        | ADHD                   | 168 subjects |          |           |           | FC from fMRI and features from sMRI, SVM, 74%                              |
| Colby et al. (2012)          | ADHD                   | HC = 491  | ADHD1 = 111, ADHD2 = 163, ADHD3 = 11, Total = 776 |           |           | FC, graph measures, ReHo from fMRI and features from sMRI, SVM, 55%        |
| Dai et al. (2012)            | ADHD                   | HC = 491  | ADHD = 285, Total = 776 |           |           | FC, ReHo from fMRI and cortical thickness, gray matter from sMRI, SVM, 67.80% |
| Dey, Rao, and Shah (2012)    | ADHD                   | HC = 307  | ADHD = 180, Total = 487 |           |           | FC between ROIs PCA-LDA, 65.60%                                           |
| Sato et al. (2012)           | ADHD                   | HC = 546  | ADHD = 383, Total = 373 |           |           | ALFF, ReHo and FC, SVM, 67%                                               |
| Fair et al. (2013)           | ADHD                   | HC = 455  | ADHD = 193, Total = 648 |           |           | FC between 160 ROIs (Dosenbach atlas), SVM, 69.20%                        |
| Wang et al. (2013)           | ADHD                   | HC = 23   | ADHD = 23, Total = 46 |           |           | ReHo, SVM, 80%                                                            |
| Anderson et al. (2014)       | ADHD                   | HC = 472  | ADHD = 476, Total = 948 |           |           | Graph measures from fMRI and features from sMRI, Decision tree, 66.80%     |
| Dey et al. (2014)            | ADHD                   | HC = 307  | ADHD = 180, Total = 487 |           |           | Graph distance measures, SVM, 73.55%                                       |
| dos Santos Siqueira et al. (2014) | ADHD               | HC = 340  | ADHD = 269, Total = 609 |           |           | Whole brain FC between 400 ROIs, SVM, 77%                                 |
| Deshpande et al. (2015)      | ADHD                   | HC = 744  | ADHD1 = 173, ADHD2 = 260, Total = 1,777 |           |           | Directional and nondirectional whole brain FC, ANN, 90%                   |
| Du, Wang, Jie, and Zhang (2016) | ADHD               | HC = 98   | ADHD = 118, Total = 216 |           |           | FC between PCA selected regions, SVM, 94.90%                              |
| Park et al. (2016)           | ADHD                   | ADHDI = 13 | ADHD2 = 21, Total = 34 |           |           | Whole brain FC between 384 ROIs during task, SVM, 91.20%                 |
| Qureshi, Oh, Min, et al. (2017) | ADHD               | HC = 67   | ADHD1 = 67, ADHD2 = 67, Total = 201 |           |           | FC from fMRI and features from sMRI, SVM, 92.90%                          |
| Riaz et al. (2018)           | ADHD                   | HC1 = 23, ADHD1 = 25; HC2 = 61, ADHD2 = 22; HC3 = 61, ADHD3 = 24; HC4 = 98, ADHD4 = 118, Total 1 = 48, Total 2 = 83, Total 3 = 85, Total 4 = 216 | |           |           | Whole brain FC between ROIs determined using the affinity propagation clustering algorithm and nonimage data, SVM, 86.70% |
et al., 2004; Mourao-Miranda, Bokde, Born, Hampel, & Stetter, 2005; Wang, Hutchinson, & Mitchell, 2004). Other recent and more efficient classifiers include: The Random Forest (Breiman, 2001), deep learning (Calhoun & Sui, 2016; Han et al., 2017; Jang et al., 2017; Plis et al., 2014; Figure 7) and artificial neural network (ANN; H. Guo et al., 2014; J. Kim et al., 2016) classifiers.

Example of multivariate machine learning methods for mental illness diagnoses includes classification of SZ using structural MRI data with classification accuracies ranging from 81% to 93% (Gould et al., 2014; Greenstein et al., 2012; Kawasaki et al., 2007; Sun et al., 2009; Yoon et al., 2008).

5.2 Multimodal studies

Although neuroimaging techniques have become popular tools to identify mental illness related biomarkers, each imaging technique has its limitations (Calhoun & Adali, 2009; Calhoun & Sui, 2016; S. Liu, Cai, et al., 2015). The modality-specific limitations can be partly overcome by developing multimodal neuroimaging techniques, by combining data obtained from multiple neuroimaging techniques, such as EEG, structural magnetic resonance imaging (sMRI), and fMRI, which provides more informative and reliable results on brain structure and dynamics than unimodal neuroimaging approach. Multimodal neuroimaging is a relatively new and rapidly expanding field that integrates data from different modalities to understand the pathophysiology of mental illness (Calhoun et al., 2006; Sui et al., 2012). For instance, by linking the genomic variation to brain function, structure and connectivity measures, imaging genomics approach can characterize the underlying neural mechanisms through which genomics variations affect cognition and behavior in mental illness. Multimodal data fusion approaches utilize complementary information available from both data modalities to jointly estimate their association by accounting for inter-modality relationships. Particularly, for complex disorders such as SZ, the relationship between features from multiple modalities may be better understood by incorporating additional information provided by advanced multimodal modeling. Thus, it is important to study the inter-relationships between multimodal neuroimaging data, as well as nonimaging features, such as, behavioral and genomic data, while performing predictive modeling of mental illness.

There are currently multiple approaches to integrate data acquired from various unimodal imaging techniques, as described by Calhoun and Sui (2016). These include a separate visual inspection of unimodal results, multimodal integration after individually analyzing unimodal results to prevent any interaction between the unimodal data, and data fusion using either asymmetric approach where one modality constrains the other (e.g., fMRI constrained EEG) or a symmetric approach where multiple modalities are analyzed jointly (e.g., joint independent component analysis [joint-ICA] or parallel independent component analysis [parallel-ICA]).

While most of the studies in the existing literature used a single modality for mental illness prediction, recent development in data fusion approaches have made multimodal neuroimaging a popular trend. Several recent studies have differentiated SZ patients from HC by combining data from rs-fMRI or task-based fMRI, and sMRI (Cabral et al., 2016; J. Ford, Shen, Makedon, Flashman, & Saykin, 2002; Qureshi, Oh, Cho, Jo, & Lee, 2017; Yang, He, & Zhong, 2016), fMRI and single nucleotide polymorphism (SNP; genomic feature; Yang et al., 2010), and rs-fMRI and MEG (Cetin et al., 2016), while only a few studies combined data from three or more modalities (Sui et al., 2013; Sui et al., 2014), with accuracy ranging from 75% to 100%. Other recent data fusion advances include integration of multiple task-based fMRI data sets (D. I. Kim et al., 2010; Sui, Adali, Pearlson, & Calhoun, 2009; Sui et al., 2015) from the same participants, where common versus specific sources of activity was specified to a greater degree than conventional general linear model-based approaches. Using a Fisher’s linear discriminate classifier, Ford and colleagues classified SZ and HC based on task-based fMRI data with 78% accuracy, and sMRI data with 52% accuracy, however, the combined multimodal data (fMRI and sMRI) resulted in the highest accuracy of 87% (J. Ford et al., 2002). Another recent multimodal neuroimaging study by Yang and colleagues integrated rs-fMRI based connectivity features and sMRI based structural features extracted using ICA, and used SVM classifier to compare unimodal versus multimodal accuracy (Yang, Chen, et al., 2016). Results from this study showed that multimodal features achieved higher accuracy (77.91%) than single modality accuracy (72.09%). Using multimodal sMRI and rs-fMRI data, Cabral and colleagues classified SZ patients and HC individuals with 75% accuracy, where the multimodal features based classification outperformed both of the unimodal features based prediction accuracy (69.7% accuracy using sMRI data and 70.5% accuracy using rs-fMRI data; Cabral et al., 2016). Qureshi and colleagues employed a similar approach to classify SZ patients and HC individuals using combined rs-fMRI and sMRI data but on a higher number of samples, and achieved 10-by-10-fold nested cross-validated prediction accuracy of 99.29% (Qureshi, Oh, Cho, et al., 2017). Note that, in order to use as much training data as possible and overcome the sample size issue, the framework utilized a nested CV without novel data for testing, which might have introduced classification bias and resulted in such high prediction accuracy. Regardless of the methodological limitations, these and other studies show the potential of leveraging multimodal imaging data. However, more robust multimodal fusion approaches and validations are required before making them available for clinical purposes.

Besides multimodal integration of MRI modalities, recent studies have also combined neuroimaging data with non-neuroimaging features, such as genomics data (e.g., SNPs), DTI, MEG, and EEG for classification of mental illnesses. Using ICA and SVM-based classifier ensemble (SVME), in a relatively small sample size and SNPs array, Yang and colleagues classified SZ patients and HC with 73.88% accuracy for SNPs data, 81.63% accuracy for voxel-level fMRI activations, 82.50% accuracy for ICA component-specific fMRI activation, and finally, 87.25% accuracy for combined fMRI-SNPs data (Honghui Yang et al., 2010). Further, in a large dataset, using multiple classifiers including sparse representation-based classifier (SRC), fuzzy c-means (FCM) classifier, and SVM-
based classifier, Cao and colleagues discriminated between SZ patients and HC individuals by combining fMRI and SNP data modalities, and found the best classification accuracy of 89.7% achieved using SRC (L. Cao et al., 2014). Another recent study by Cetin and colleagues integrated rs-fMRI and MEG data to distinguish SZ from HC, and found that the best performance of 87.91% accuracy was obtained by using the ensemble classifier (Cetin et al., 2016; Figure 8). Using a novel data fusion technique known as mCCA + jICA and multiple types of classifiers, Sui and colleagues integrated features from rs-fMRI, sMRI, and DTI (i.e., FA) to classify SZ patients and HC individuals, and achieved a maximum classification accuracy of over 90% using radial basis function support vector machine (RSVM) classifier on DTI (i.e., FA) and sMRI (gray matter) features (Sui, He, Pearlson, et al., 2013). Further, using the sample data fusion technique (i.e., mCCA+jICA) and features from rs-fMRI, sMRI, and EEG modalities, Sui and colleagues utilized a SVM classifier with recursive feature elimination (SVM-RFE), and obtained 91% accuracy in training data and 100% accuracy (i.e., prediction rate) with combination of all modalities to classify SZ patients and HC individuals (Sui et al., 2014). In order to classify ultra-high-risk individuals for psychosis, first-episode psychosis and HC, Pettersson-Yeo and colleagues used a multi-step data fusion approach that includes an unweighted sum of kernels, multi-kernel learning, prediction averaging, and majority voting, and obtained 86.33% accuracy by combining features from DTI and fMRI modalities (Pettersson-Yeo et al., 2014).

The results of the above-mentioned studies are encouraging for using multimodal neuroimaging in classification of mental illness, suggesting that data fusion methods combined with advanced machine learning techniques present a promising direction for mental illness prediction.

### 5.3 Multi-class classification for disease subtype to reduce diagnoses heterogeneity

While the case (i.e., patient with mental illness) versus control diagnostic approach has been successfully implemented in the existing mental illness prediction literature, it does not address the differential diagnosis aspect of mental illness prediction (i.e., distinguishing between illnesses with overlapping symptoms or subgroup diagnosis). Traditional case versus control model ignores the heterogeneity within the disease group by assigning predefined case or control label to the test sample. Many severe mental illnesses, such as SZ, schizoaffective disorder, unipolar and bipolar depression, and other mood disorders share substantial overlapping symptoms, and therefore disease prediction based on the reported symptoms alone is not adequate for accurate diagnosis. To address this, the National Institute of Mental Health introduced the research domain criteria (RDoC; http://www.nimh.nih.gov/research-funding/rdoc), as a way to classify mental illnesses based on multiple symptom, behavioral and biological dimensions, with a goal of reducing heterogeneity across diagnostic groups (T. Insel et al., 2010; T. R. Insel, 2014). Consistent with the European roadmap for mental health research (Schumann et al., 2014), the RDoC offers an improved classification validation at the neurobiological level. Most of the studies addressing the RDoC initiative have implemented various data-driven, clustering approaches based on neuropsychological measures in order to divide the clinical populations. These studies include subtype diagnoses of SZ (Bell, Corbera, Johannesen, Fiszdon, & Wexler, 2011; Brodersen et al., 2014), mood disorders (Lamers et al., 2010; Van Loo, De Jonge, Romeijn, Kessler, & Schoevers, 2012), and ADHD (Dias et al., 2015; D. A. Fair, Bathula, Nigg, & Nigg, 2012; Koutsouleris et al., 2015; van Hulst, De Zeeuw, & Durston, 2015).

As discussed above, one of the main limitations of the traditional case-control prediction is the binary disease characterization, where test samples are assigned to either case or control category. This approach overlooks the associated disease heterogeneity, commonly known as the disease subtype. However, many heterogeneous mental illnesses including autism and SZ are defined as spectrum disorders (i.e., a continuum) with multiple disease etiologies lying under the same diagnostic category. While it is a common practice to classify such spectrum-like disorders using the generic category to find diagnostic biomarkers, a major issue in mental illness diagnostic procedure is the lack of differential diagnosis of patients across several disease subtypes. Accurate diagnosis of disease subtype is very critical for the appropriate course of treatments. For example, in case of SZ, patients can exhibit similar cognitive deficits but with variable magnitude. Therefore, to emphasize the phenotypic heterogeneity in SZ, two major subtypes with different genetic and cognitive profiles have been introduced: (a) cognitive deficit and (b) cognitively spared (Green et al., 2013; Jablensky, 2006). However, differential diagnosis of SZ subtypes has been rarely studied, due to limited sample size. The number of subjects in each disease subtype is small in most of the existing datasets, which limits the ability to develop robust subtype predictor to accurately differentiate them. Ingalhalikar and colleagues proposed an unsupervised spectral clustering approach using multi-edge graphs derived from a structural connectivity network among 78 ROIs to classify subtypes of autism and SZ (Ingalhalikar et al., 2012).

Among the surveyed studies in this review, only a few considered the important area of automatic differential diagnosis. Costafreda and colleagues used fMRI with a verbal fluency task for subject-level classification of SZ, bipolar and HCs (Costafreda et al., 2011). Two studies by Calhoun et al. (2008) (Figure 9) and Arribas et al. (2010) used fMRI with an auditory oddball task where they have applied an ICA approach to extract the default model network related features and the temporal lobe of the brain. Both of these studies achieved high prediction accuracy between SZ and BP disorder. Rashid and colleagues proposed static dynamic functional network connectivity (FNC) features-based algorithms (Figure 10) to automatically classify SZ, bipolar and HCs (Rashid et al., 2016). Another study by Pardo and colleagues used a combination of volumes of 23 ROIs derived from structural MRI along with 22 neuropsychological test scores for automatic classification of SZ, bipolar and HCs (Pardo et al., 2006). More recently, using gray matter densities, Schnack and colleagues
proposed a classification framework for SZ, bipolar and HCs (Schnack et al., 2014). Using gray matter maps from structural MRI, Koutsouleris and colleagues classified SZ from mood disorder (Koutsouleris et al., 2015). Ota and colleagues combined volumetric measures derived from structural MRI with FA from dMRI in selected ROIs to classify SZ from MDD (Ota et al., 2013). Moreover, using gray matter volumes of caudate and ventral diencephalon, Sacchet, Livermore, et al. (2015) proposed an algorithm to classify MDD, bipolar and remitted MDD patients.

Although the limited sample size in most of the current datasets makes it challenging to perform disease subtype prediction as the number of subjects in each diseases subtype is small, it shows the potential for a paradigm shift in the predictive modeling of spectrum-like mental illness beyond discrete, case–control diagnosis.

### 5.4 Advanced algorithms for brain-based prediction

Recent advanced machine learning algorithms have shown tremendous potential for neuroimaging-based mental illness prediction. For example, a recent study proposed a novel parallel group ICA-based framework to jointly estimate the association between functional network variability and structural covariation in SZ, as well as to predict several cognitive domain scores based on these associated functional/structural features (Qi et al., 2019; Figure 11). Briefly, by jointly incorporating and estimating temporal domain features from fMRI (extracted from group ICA) and structural MRI features within a parallel group ICA algorithm, functional network variability and structural covariation were jointly estimated to identify between-modality linkage. Using real neuroimaging data, a significant functional and structural MRI component pair was identified that captured group difference in both imaging modalities, which further correlated with cognitive scores suggesting that multimodal brain features can predict multiple cognitive scores. Another recent SZ study proposed a multimodal fusion with reference algorithm by combining multi-site canonical correlation analysis with reference and joint-ICA (MCCAR+jICA) to identify co-varying multimodal feature patterns using a reference (specifically, working memory performance) in a three-way data fusion (fMRI, sMRI, and dMRI; Qi et al., 2018). Results identified several brain regions that were previously linked with working memory deficits in SZ, suggesting that the novel MCCAR+jICA method has great potentials to identify biomarkers for severe mental disorders, such as SZ. Further, Sui and colleagues implemented a constrained fusion approach to predict cognition in SZ (Sui et al., 2015; Figure 12). The assessment of cognition was measured using the MATRICS Consensus Cognitive Battery (MCCB), and using multi-set canonical correlation, the linkage between MCCB and brain abnormalities as measured by fractional amplitude of low-frequency fluctuations (fALFF) from resting fMRI, gray matter density (GM) from structural MRI, and FA from dMRI were explored. Findings from this study suggested that the associated functional and structural deficits might be linked to cognitive impairments in SZ. Other recent neuroimaging-based data-driven advanced algorithms include biclustering and triclustering ICA approaches that utilize spatial and temporal variance as a measure to cluster mental disorders into homogeneous subgroups. For instance, using gray matter concentration (GMC) from SZ patients, Gupta and colleagues implemented source-based morphometry (SBM) decomposition, followed by subtype component reconstruction using group information-guided ICA (GIG-ICA), and identified two subtypes (i.e., two different subsets of subjects; Gupta et al., 2017). Also, Rahman and colleagues have used structural MRI features from SZ patients to perform multi-component and symptom bi-clustering (Rahaman et al., 2019), and further extended this approach into tri-clustering framework using dynamic FNC measures to identify disease subtypes (Rahaman, Damaraju, & Calhoun, 2019).

### 5.5 Functional connectivity measures for brain-based prediction

In recent years, brain connectivity studies using neuroimaging have become popular to investigate the associations among brain networks. Functional connectivity (FC) can be quantified using a variety of different neuroimaging techniques. A commonly used measure is fMRI, which measures synchronized brain activity via blood oxygenation and infers functional interactions among different brain regions (Craddock et al., 2013). FC, defined as temporal correlation (or other types of statistical dependency) among spatially distant brain regions (K. Friston, 2002), has recently been used to examine the functional organization and temporal dependencies among these remote brain regions. Different analytic tools have been applied to resting-state fMRI data to describe brain functional connectivity. Two widely used FC approaches are (a) seed-based analysis (B. Biswal, Zerrin Yetkin, Haughton, & Hyde, 1995; Greicius, Krasnow, Reiss, & Menon, 2003) and (b) purely data-driven methods, such as ICA (Calhoun & Adali, 2012; Calhoun, Adali, Pearlson, & Pekar, 2001a; Calhoun, Eichele, & Pearlson, 2009; Fox & Raichle, 2007). FC can also be investigated at the network level using spatial ICA, and connectivity among spatial components is referred to as FNC (Jafari, Pearlson, Stevens, & Calhoun, 2008). In most common cases, FC (as well as FNC) is considered to be stationary over the entire scanning period, formally known as the sFC/sFNC analysis.

For predictive analyses, specific methods for measuring functional connectivity might influence the features of interest as well as the classification accuracy. Given the high dimensional features resulting from FC analysis, without efficient but meaningful feature selection, the classifier could introduce overfitting problem, leading to poor classification performance. Thus, to eliminate redundant features and choose only appropriate features from FC measures, application of suitable feature selection strategies is critical. For FC based feature selection, approaches such as filter (e.g., statistical test, Fisher score and correlation coefficient; Figure 13; J. S. Anderson et al., 2011; Arbabshirani et al., 2013; Bassett et al., 2012; Calhoun et al., 2008; W. Du et al., 2012; Guyon & Elisseeff, 2003), wrapper (e.g., recursive feature elimination [RFE], sequential feature selection and genetic
algorithm; Fan et al., 2011; Guyon & Elisseeff, 2003; Venkataraman et al., 2012; Yu, Shen, Zhang, et al., 2013), and embedded (e.g., LASSO regularization, decision tree; Fonti & Beltser, 2017; B. Jie et al., 2014; Lal, Chapelle, Weston, & Elisseeff, 2006; Watanabe et al., 2014) methods can be applied.

Table 9 presents FNC-based classification studies for automatic diagnoses of SZ, MDD/BP, ASD, and ADHD patients. Both FC and FNC measures have been utilized for subject-level diagnosis (i.e., prediction) of severe mental disorders, such as SZ and bipolar. For example, by using the connectivity between functional brain networks (i.e., FNC) identified by ICA several recent studies have classified SZ (80% accuracy (A. Anderson & Cohen, 2013) 63–96% accuracy (Arbabshirani et al., 2013); 84.4% accuracy (Kaufmann et al., 2015). Another popular estimate for FC between ROIs identified based on different atlases has also been used to distinguish between SZ and healthy individuals (75% accuracy (Venkataraman et al., 2012); 82.8% accuracy (Su et al., 2013); 80.9% accuracy (Yu, Shen, Zeng, et al., 2013), 62% accuracy (Yu, Shen, Zhang, et al., 2013); 59.7–90.8% accuracy (Watanabe et al., 2014); 86% accuracy (J. Kim et al., 2016)). Further, Bassett et al. (2012) have use high-level network organization as features for classification of SZ (75% accuracy). In addition to the single modality based FNC features, studies have also explored the fusion of FC with features from other modalities to classify SZ. For example, Yang and colleagues developed a hybrid machine learning approach using fMRI (voxels and ICA networks) and SNP features for classification of SZ and achieved 87% classification accuracy using the combined features (i.e., voxel, ICA network, and SNP data; Yang et al., 2010). Recent studies have also classified SZ and BP disorder using FC-based features. Using ICA spatial maps of the temporal lobe and the default mode networks, Calhoun and colleagues classified SZ and bipolar patients with accuracies of 83–95% (Calhoun et al., 2008). Another study used the ICA spatial maps as features and a probabilistic Bayesian classifier to discriminate between SZ, BP disorder, and healthy individuals, and achieved the average three-way correct classification rate within the range of 70–72% (Arribas et al., 2010). Few recent studies also performed of classification of mental illnesses with overlapping symptoms, such as SZ, schizoaffective disorder, and BP disorder with psychosis (Cardno & Owen, 2014; Cosgrove & Suppes, 2013; Pearson et al., 2016). Du and colleagues used GIG-ICA to extract resting-state brain networks and classified SZ patients, BP disorder with psychosis, schizoaffective disorder with manic episode disorders schizoaffective disorder with depressive episodes exclusively and healthy individuals (Du et al., 2015). FNC features were selected using RFE method and a five-class SVM classifier was used for training purpose, achieving 68.75% of classification accuracy (Figure 14).

FC measures have also been utilized as features for classification of autism and ADHD. Using resting-state FC measures from 7,266 ROI unclosing only the gray matter, Anderson and colleagues achieved 89% accuracy for the subjects <20 years age and at 79% for all subject (J. S. Anderson et al., 2011). Murdaugh and colleagues used both

![FIGURE 14](image-url) An example showing prediction of schizophrenia and bipolar using network measures and hierarchical clustering. (a) Distance matrix from the feature vectors. (b) The mean inter-group and intra-group distance matrix. (c) The Results from t-distributed stochastic neighbor embedding (t-SNE) method showing the projection results of subjects, where each point refers to a subject (group-wise colored). (d) The linkage results from the hierarchical clustering method. BP, bipolar; HC, healthy control; SADD: schizoaffective (depression); SADM, schizoaffective (manic); SZ, schizophrenia. Figure reprinted with permission from Du et al. (2015)
seed-based FC and whole-brain FC in a logistic regression classifier to classify autism and reported 96.3% accuracy with both whole-brain and seed-based FC features (Murdaugh et al., 2012). Further, using FC between three ROI sets from ABIDE dataset, Plitt and colleagues applied RFE-based feature selection and both logistic regression and SVM classifiers to classify autism and achieved an overall 76.7% accuracy (Plitt et al., 2015). In a multimodal classification study of autism, Deshpande and colleagues used FC estimates and fractional anisotropy (from DTI data) and obtained a maximum classification accuracy of 95.9% with a recursive cluster elimination based SVM classifier (Deshpande et al., 2013). Another recent resting-state FC-based (between 90 ROIs) prediction study used deep learning classifier (probabilistic neural network [PNN]) for classification of ASD and achieved a classification accuracy of about 90% (Iidaka, 2015). Interestingly, FC between signals in different frequency bands was used as features in a recent ASD classification study, where the Slow-4 band (0.027–0.073 Hz) was found to capture the most discriminative features (H. Chen et al., 2016). To discriminate ADHD from healthy individuals, Zhu and colleagues used a PCA-based Fisher discriminative analysis (PC-FDA) with regional homogeneity (ReHo) from fMRI data as features, and showed a classification accuracy of 85% (C.-Z. Zhu et al., 2008). Another study by Wang and colleagues also used ReHo from resting-state fMRI data in a SVM classifier, and obtained a classification accuracy of 80% for discriminating ADHD from healthy individuals (Wang et al., 2013). Several other studies also used FC measures to successfully classify ADHD from HCs (Dey et al., 2014; D. Fair et al., 2013; João Ricardo Sato et al., 2012). Moreover, by leveraging a large-scale resting-state fMRI study of SZ from multiple sites (i.e., human connectome project), another recent study proposed a ICA-based preprocessing pipeline to extract FNC and spatial map based imaging features as potential biomarkers. Results showed that compared to FNC-based features, spatial map shows better classification performance in all experiments (Lin et al., 2019). Another study proposed a robust group information guided ICA (GIG-ICA) based framework to estimate functional network maps and connectivity for linking neuromarkers among different diseases and separate studies, where the connectivity measures were independently computed and optimized to achieve independence based on each coming individual-subject data. Findings from this study showed that the network features computed using the proposed method were more effective for predicting different diseases and classifying patients (Y. Du et al., 2019). One of the advantages of this framework is it does not require selecting components of interest and can be fully automated (Figure 15).

![Figure 15](image-url)
COMMON SHORTCOMINGS IN PREDICTOME STUDIES

6.1 Feature selection and reduction

An additional limitation of mental illness prediction studies includes the selection and dimensionality reduction of meaningful features. Almost all of the papers surveyed in this review have group-level discriminative analysis followed by subject-level classification. Many of these studies have first performed discriminative analyses using statistical tests (e.g., t tests) to extract significant features showing group difference, and then using these features performed subject-level classification. However, the use of test dataset together with the training dataset during feature selection, extraction or reduction will introduce additional bias to the predictive model. This process of feature selection based on the group differences results identified from the whole sample could cause a "double dipping" issue, that may lead to a biased performance (Bishop, 2006; Demirci, Clark, Magnotta, et al., 2008). Another major issue with the group difference based feature selection approach is that the significance levels are based on
between errors on training data set and margin maximization. A defined soft margin hyperparameter that affects the trade-off in high-dimensional space. Linear SVM classifier includes a user-binary classification that maximizes the boundary between classes in stage, typically during the CV of the training samples. One of the hyperparameter optimization step is performed during the training in order to achieve the best performance of the algorithm. The choosing a set of optimal hyperparameters for a learning algorithm hyperparameter optimization or tuning, refers to the problem of optimizing for RBF kernel and degree for polynomial kernel). Therefore, inefficient hyperparameter optimization could negatively influence model performance.

6.2 Overfitting

Overfitting refers to a model that models the training data too well, resulting in very good classification performance on the training data (i.e., observed data) but very poor performance on independent, testing data (Pereira et al., 2009). Overfitting can be caused by utilizing models with large number of features from small sample size or complex models with many parameters, as the model would capture noisy features from the data more than the actual features of interest (Franke et al., 2010; Klöppel et al., 2008). Since neuroimaging datasets on mental illnesses have smaller samples in general and many features of interest, predictive models using these datasets are susceptible to overfitting. The majority of the surveyed studies reported in the current work performed predictive modeling based on a very small number of subjects, resulting in a decrease in the overall classification accuracy with smaller sample size. CV and regularization are common approaches to control for overfitting in neuroimaging data-based predictive modeling.

6.3 Optimal model selection

In predictive modeling, model selection, more formally known as hyperparameter optimization or tuning, refers to the problem of choosing a set of optimal hyperparameters for a learning algorithm in order to achieve the best performance of the algorithm. The hyperparameter optimization step is performed during the training stage, typically during the CV of the training samples. One of the most commonly used classifiers is the SVM which is designed for binary classification that maximizes the boundary between classes in a high-dimensional space. Linear SVM classifier includes a user-defined soft margin hyperparameter that affects the trade-off between errors on training data set and margin maximization. A smaller soft margin would introduce more errors, resulting in a larger margin. Nonlinear SVM classifier includes additional hyperparameters depending on the kernel of choice (e.g., sigma/gamma for RBF kernel and degree for polynomial kernel). Therefore, inefficient hyperparameter optimization could negatively influence model performance.

6.4 Challenge with reproducibility

In the existing brain-based prediction literature, variability across raw data processing and analysis streams, feature types, feature selection schemes, choice of classifier, and CV methods may limit reproducibility of outcomes across independent datasets. Indeed, without any standard approach, the growing flexibility across machine learning pipelines is introducing less reliable replication across studies (Squeglia et al., 2016). In order to provide optimum diagnostic tools, the predictive models should be able to handle new samples obtained from independent research facilities. Further, the models should perform consistently across different data-acquisition devices (e.g., Siemens/Philips/GE scanners), across different geographic areas (e.g., United States, Europe, etc.), and across diverse populations (e.g., same mental illnesses with distinctive comorbidity). Also, the performance should persist across different evaluation metrics (e.g., accuracy, sensitivity, specificity, etc.), as well as different medical settings (e.g., rural vs. city hospital; Woo, Chang, Lindquist, & Wager, 2017).

6.5 Prediction outcome comparison across studies

A common practice in prediction studies is to highlight the overall accuracy as the ultimate model performance measure. While many surveyed studies have reported that their algorithms performed better than other existing studies, this claim is often unsubstantiated and solely based on the comparison between their overall accuracy and that of the existing studies. Without well-matched study variables (i.e., sample size, age, sex, scanner parameters and imaging sequences, medication, symptom scores, data modality, length of scanning for functional data, preprocessing pipeline, features of interest, feature selection methods, classifier type, and CV scheme), these comparisons are simply impractical. Further, any such performance-related claims must be made after appropriate statistical significance tests.

6.6 Heterogeneity between patients

Another limitation for neuroimaging based machine learning studies is the substantial heterogeneity that exists between patients. In research-based neuroimaging studies, participants are recruited based on well-matched age, sex, or education background, typically with a particular type of brain pathology. In contrast, participants recruited in...
the clinical settings may include several types of pathology with variability in disease stage and demographic variables (i.e., age, sex etc.). As mentioned earlier, classification performance (e.g., accuracy) can be improved by using a larger training sample (Franke et al., 2010; Klöppel et al., 2008), and may also reduce the disease heterogeneity by integrating the whole spectrum of clinical and pathological features. Recent approaches, such as outlier detection (i.e., treating patient classification as an outlier detection problem (Mourão-Miranda et al., 2011) can be implemented to reduce higher heterogeneity across patients. Further, two recent studies have performed multi-component and symptom bi-clustering where homogeneous clusters along various dimensions were identified for SZ subtype prediction (Gupta et al., 2017; Rahaman, Turner, et al., 2019; Figure 16).

7  |  FUTURE DIRECTIONS FOR BRAIN-BASED PREDICTOME STUDIES

7.1  |  Prediction using deep learning techniques

In recent years, deep learning, which is a family of machine learning techniques, has gained significant attention for patterns recognition and classification in various data analysis fields including neuroimaging. The fundamental difference between traditional machine learning and deep learning is that, deep learning has the capability to detect and learn complex patterns from the raw data by utilizing consecutive nonlinear transformations. Due to its ability to identify complex and subtle (and potentially nonlinear) patterns based on data-driven automatic feature learning process, deep learning has recently become an attractive tool in neuroimaging studies of mental illness. Deep learning-based approaches are more effective when examining large number of features without any proper knowledge in feature selection. Using a hierarchical model of nonlinear layers, deep learning can model more complex data patterns. To summarize, the main advantages to use deep learning over traditional machine learning approaches include: (a) ability to implement data-driven automatic feature learning and eliminating the subjectivity related to feature selection, and (b) the depth of model design that includes a hierarchy of nonlinear layers, allowing the model to efficiently identify complex data patterns.

Deep learning offers a promising tool for understanding the neural basis of mental illness by extracting complex, hidden patterns from high-dimensional neuroimaging data (Kriegeskorte, 2015). Deep models have multiple advantages, including requiring exponentially smaller number of parameters to model the same thing as models by traditional machine learning approaches (Bengio, 2012). Further, for larger neuroimaging datasets, deep learning provides the opportunity to more efficiently diagnose mental illnesses (E. Castro, Ulloa, Plis, Turner, & Calhoun, 2015). In order to examine mental illness using deep learning, several studies have utilized a wide range of

FIGURE 17  An imaging-genomics framework to jointly estimate group differences in schizophrenia (SZ) and healthy controls (HC). The parallel ICA based multimodal framework incorporating imaging (dynamic FNC measures) and genomic (single nucleotide polymorphism, SNP) data. Figures reprinted with permission from Rashid et al. (2019)
neuroimaging modalities including sMRI, resting fMRI, and multimodal data. For instance, using deep belief networks on both structural and functional MRI data, Plis and colleagues have demonstrated that deep learning methods are able to generate physiologically meaningful MRI-based features that can uncover associations from high-dimensional MRI data (Plis et al., 2014). Another study by Hjelm and colleagues used restricted Boltzmann machines (RBM) to capture intrinsic brain networks in fMRI data showed that RBMs could extract spatial networks and their activation with the accuracy of traditional matrix factorization methods such as ICA (Hjelm et al., 2014). Note that, RBM can be stacked to obtain deeper models as needed for deep learning which is a limitation in traditional approaches such as ICA, non-negative matrix factorization (NMF), or sparse PCA. More recently, deep learning has been employed in classification of patients using neuroimaging data. Kim and colleagues used deep learning for classification of SZ patients from HCs based on functional connectivity patterns, and showed that their approach outperforms SVM by a significant margin (J. Kim et al., 2016). Using structural MRI data, Castro and colleagues developed a nonlinear independent component estimation (NICE) which is an extension of ICA using deep architecture, and detected gray matter abnormalities in patients with SZ. The resulting NICE components captured decreased patterns of GMC in SZ, showing potential for application in SZ prediction studies (Castro et al., 2016). Further, Jang and colleagues applied fully connected feed-forward deep neural networks (DNN) to four sensorimotor task-based fMRI data from 12 HC individuals. More specifically, the framework used a restricted Boltzmann machine-based deep belief network classifier and obtained lower minimum error rates (mean ± SD; three-layer DNN = 6.9 ± 3.8%, single-layer DNN = 9.4 ± 4.6 and two-layer DNN = 7.4 ± 4.1). Results showed that the DNN is able to classify a single-volume fMRI extracted by hidden representations of task-specific fMRI volumes distributed across multiple hidden layers, showing great potential for automatic mental illness prediction using fMRI volumes in cases where limited estimates of activation patterns or ad hoc statistical evaluation are available (Jang et al., 2017). Plis and colleagues implemented a deep learning method to identify association between multimodal neuroimaging data (i.e., sMRI and fMRI) in 144 SZ patients and 154 HC individuals. Dynamic FNC states and ICA-based gray matter density maps were used as functional and structural features, respectively. Using the hidden association between brain structure and function, this study distinguished between SZ and HC participants (Plis et al., 2018). Another multimodal data integration study developed a neural network framework to study the brain development. Briefly, this study proposed a deep collaborative learning (DCL) method to investigated the difference of functional connectivity measures across different age groups, and achieved a maximum prediction accuracy of about 98% (W. Hu, Cai, Zhang, Calhoun, & Wang, 2019).

Within the field of neuroimaging, deep learning techniques have shown to achieve superior performance over many conventional machine learning approaches; however, currently there are some limitations and challenges of deep learning techniques which require further improvement prior to practical implementations. Some of the limitations of deep learning are discussed as follows:

1. Deep learning techniques are highly dependent on the quality and amount of training data which may result into overfitting (i.e., learning unrelated variations in the data), which may, in turn, result in lack of generalizability. Approaches such as regularization strategies (e.g., L1 or L2 norms, dropout, and weight decays) are being incorporated in deep learning methods in order to address the issue of overfitting (Hosseini-Asl, Gimel’farb, & El-Baz, 2016; J. Kim et al., 2016; F. Li, Huang, et al., 2014; S. Liu et al., 2014). The regularization strategies can improve the performance of the deep learning algorithms. Other approaches to avoid overfitting include dimensionality reduction (e.g., using region-level features instead of voxel-level features, selecting features using principal component analysis) of the data prior to training them using the model. Although, current evidence suggests that most deep learning models, including convolution neural network (CNN), can perform well with voxel-level neuroimaging data without feature selection or reduction (Hosseini-Asl et al., 2016).

2. Deep learning techniques require a large amount of training data to identify more generalized features and improved performance, and therefore demonstrate poorer performance with small sample size (Nie, Zhang, Adeli, Liu, & Shen, 2016). Further, noisy neuroimaging data may not improve the performance of the model beyond linear methods even with increasing sample size (e.g., ImageNet). One solution to this issue could be to use specific types of less-noisy features to train the model. For example, in neuroimaging, voxel-level features tend to be very detailed and noisy, whereas region-level features show less sensitivity to noise as they overlook more localized patterns. Further, the accuracy of deep learning techniques may be improved by making full use of the automatically extracted features even for studies with smaller sample size.

3. Deep learning models offer a black box-like system, which may introduce lack of transparency during the learning and testing steps (Alain & Bengio, 2016; Yosinski, Clune, Nguyen, Fuchs, & Lipson, 2015). In many cases, it is very difficult to understand the technical and logical bases of the model. The lack of transparency of deep learning may limit the interpretability of the neuroimaging results. Indeed, the multiple nonlinearities within deep models make it challenging to trace back the successive layers of weights to the original brain data, therefore limiting the ability to detect abnormalities within brain regions (Suk et al., 2015). To address this issue, currently two approaches are being implemented: (a) input modification methods (i.e., visualization techniques by systematically modifying the input, the resulting changes in the output, and in intermediate layers; example: occlusion method; Zeiler & Fergus, 2014) and (b) deconvolution methods (i.e., determining the contribution of features of the input to the output data; example: guided back-propagation; Springenberg, Dosovitskiy, Brox, & Riedmiller, 2014).

4. Another challenging issue for deep learning methods is the workflow integration, particularly for clinical use of these models.
In order to implement these deep models in the clinical settings, it is necessary to share the relevant knowledge of these models with the clinicians, as well as receive feedback from them as the end-users during the model development stage.

7.2 | Large-scale/big data approaches

The most commonly observed limitation in the field of mental illness predictive studies is the limited sample size. The lack of larger datasets (i.e., number of subjects) as well as insufficient phenotypic details, such as clinical record, comorbidities, symptom and disease progression, treatment outcomes, and responses limit the scope of utilizing machine learning algorithm for developing personalized care of mental illness. The majority of the prediction studies in this review have a relatively small sample size. Although sample size is a common limitation across all prediction studies, compared to the other fields of studies in which machine learning is used, neuroimaging studies in general have very small sample size due to the difficulties of recruiting patients and the cost of collecting the data and the data also have high dimensionality. This may introduce several issues including decrease in classifier performance (Franke et al., 2010; Klöppel et al., 2009), lack of generalization for diagnosis purposes, and inability to address disease heterogeneity, and model overfitting due to poor sample size (Pereira et al., 2009). For optimal evaluation of machine learning methods, therefore larger sample size is required to minimize the variance in assessments of accuracy, sensitivity, specificity, and other performance measures.

To address this limitation within neuroimaging research, multiple ongoing efforts have created dataset repositories. The large-scale or “big data” revolution shows promises to reduce data heterogeneity related issues in neuroimaging studies (Franke et al., 2010; Klöppel et al., 2008). While most existing neuroimaging studies consist of moderate sample sizes, typically less than 50 subjects, a few number of studies have started to embrace the “big data” for neuroimaging by either collecting tens of thousands of subjects or taking advantage of huge increases in the quantity of imaging and nonimaging features collected on each subject. Currently, there exist three types of big-data initiatives: (a) centralized, (b) decentralized data sharing repositories, and (c) large-scale studies.

7.2.1 | Centralized repositories

Centralized data sharing repositories offer open sharing of data. Many of these repositories include multiple sites and commercial cloud computing setups to provide support with high computational load. Some examples of centralized repositories (there are many dozens of these currently) include:

1 NIMH data archive: The Adolescent Brain Cognitive Development Study (ABCD) (Casey et al., 2018).
2 NITRC: For SZ, ASD, and ADHD (Buccigrossi et al., 2008).
3 COINS: Collaborative Informatics and Neuroimaging Suite or COINS for ASD patients (https://coins.mrn.org/).

Note that, these are only some examples of what have been shared via the centralized repository platforms.

7.2.2 | Decentralized repositories

Additionally, to address the legal, ethical, and sociological concerns that might prohibit open data sharing initiatives and avoid re-identification of study subjects, repositories with anonymized raw data are also being established.

1 COINTAC: a decentralized and privacy-enabled infrastructure model for brain imaging data (Gazula et al., 2018; Plis et al., 2016).

Further information on implementation of decentralized algorithms, enhancement of user interface, regression statistic calculation for decentralization, and comprehensive pipeline specifications can be found in Ming et al. (2017).

7.2.3 | Large-scale studies

Moreover, several multi-site studies have also started to share neuroimaging data in a collaborative setup. These studies include:

1 FBIRN: For SZ patients (J. M. Ford et al., 2008).
2 COBRE: For SZ patients (center of biomedical research excellence, http://cobre.mrn.org/).
3 MCIC: For SZ patients (Gollub et al., 2013).
4 Functional Connectomes Project: For healthy subjects (B. B. Biswal et al., 2010).
5 ABIDE: For ASD patients (Di Martino et al., 2014).
6 ADHD-200: For children with ADHD (Consortium & others, 2012).

While the data sharing initiatives show potential (Milham et al., 2018), there are several methodological challenges of big data approaches in the field of neuroimaging which need to be addressed.

1 The lack of uniformity in data acquisition and processing across contributing sites could introduce performance bias of predictive modeling while using the pooled multi-site data. For large-scale imaging studies, there are several potential artifacts of concern which may include factors affecting imaging and other variables of interest, such as (a) head motion, (b) head size, (c) heart rate and respiratory variations, (d) variability in scanner hardware, and (e) variability in scanner hardware. To mitigate the effects of these confounding factors, more sophisticated image processing techniques should be developed (for examples of such processing techniques, see (1) Salimi-Khorshidi et al., 2014); rsfMRI data, (2) Andersson & Sotiropoulos, 2016; dMRI data, and (3) Fortin
et al., 2017; Fortin et al., 2018; Johnson, Li, & Rabinovic, 2007); dMRI and sMRI data. Further efforts are required to standardize data acquisition parameters across all data sharing sites for more ideal data pooling. For example, when sMRI data were analyzed, inconsistency in field strength and imaging sequence design showed evidence of significant systematic differences in multi-site studies (Fennema-Notestine et al., 2007; Stonnington et al., 2008). Without standardized parameter agreement, the variability observed across the subjects is increasingly driven by variability in scanner and imaging parameters, which could potentially introduce false disease-specific effects. Another study showed similar field related differences in 10,000 subjects across 1.5 T and 3 T scanners, they also showed evidence of highly consistent changes associated with age (Panta et al., 2016). By designing and maintaining study protocols across all contribution sites, and developing analytic approaches which are more robust to site effects (e.g., end-to-end deep learning to predict and remove site effects), chances of observing disease-specific effects are increased. The benefits of big data are many, and with improved integration between participating sites, in term of acquisition and other parameters, the potential inhomogeneities can be mitigated.

2 Another major issue for “big” imaging data is the statistical challenge. These rich datasets are designed to explore a variety of hypotheses. As researchers investigate multiple imaging modalities, many of them tend to explore various alternative models to search for significance without proper multiple comparison testing or CV framework. This makes CV and replication even more essential. Similarly, the effect size should also be reported, if using null hypothesis testing, big data can provide highly significant results for tiny effect sizes, which may not be particularly useful for any individual subjects. Also, the use of robust test statistics is important, for example, the use of, nonparametric testing, such as permutation-based tests can be incorporated while examining multiple modalities (Winkler et al., 2016).

3 With big data comes the “curse of dimensionality”. Compared to the number of observations, there are many features in high dimensional data, making it susceptible to issues such as sparsity, multicollinearity, computational cost, model complexity, and overfitting. One potential solution could be to implement feature selection or reduction approaches, such as principal component analysis, prior to analyzing and modeling the data.

4 Neuroimaging data sharing through the big data consortiums have raised some ethical and privacy concerns, for example, the possibility of facial reconstruction from structural images. By removing recognizable facial features using the de-facing approach prior to data sharing, this concern can potentially be addressed. Other ethical concerns include the risk of subject identification based on their geographical location, since typically these large-scale studies are conducted within a particular region. By adopting multilayer, restricted data sharing approach, more controlled access to the full dataset can be achieved, thus eliminating the risk of subject identification. Another approach is to utilize federated learning or decentralized analysis approaches, for example, the COINSTAC tool allows one to perform regression as well as more advanced voxel-wise and machine learning-based approaches in a decentralized framework without requiring the data to be shared (Plis et al., 2016).

7.3 | Standard machine learning competitions in neuroimaging

In the field of machine learning, recent standard predictive analyses-based competitions have already contributed to the growth of technologies. Typical layouts of such competitions include: (a) provide the participants with a labeled training dataset and an unlabeled testing dataset, (b) based on the training dataset, the participants attempt to develop a predictive model with best performance, and then apply the trained model to predict the testing samples’ labels, and (c) finally, the participants submit the prediction results and performance. Using the standard training and testing dataset and some basic preprocessing provided by these competitions, the participants can focus on the predictive modeling aspects without any biased outcomes. Although, in neuroimaging, such competitions are not as common as other fields due to the restricted data sharing policies, a number of brain-based machine learning competitions have been held in recent years. In 2011, the ADHD-200 competition was held which included resting-state fMRI data, as well as anatomical and phnototypical data of 776 subjects (491 typically developing children and 285 ADHD) for training purposes and 197 subjects as testing dataset (Consortium & others, 2012), with the goal of distinguishing between patients with ADHD and healthy, typically developing children. This competition demonstrated a greater prospect toward multi-site, large-scale ADHD data sharing. Another more recent machine learning competition was organized by the IEEE MLSP workshop that provided only neuroimaging-based features, with the goal of automatic prediction of SZ patients from HCs (Silva et al., 2014). Briefly, the participants were provided with FNC measures from resting-state fMRI, as well as the ICA loadings of SBM measures from structural MRI from 144 subjects (75 HCs and 69 SZ patients). The competition included 245 participating teams, with the winning team obtaining an AUC of around 0.90. Additionally, an AUC of around 0.94 was achieved by combining the top three models (Silva et al., 2014). Further, in 2018, to challenge participants in predicting MDD and HCs using structural MRI data, the predictive analytics competition (PAC) was arranged (https://www.photon-ai.com/pac). The competition included training data from 759 MDD patients and 1,033 HCs and unlabeled testing data from 448 subjects obtained across three different publicly available sites. The winners achieved a classification accuracy of 65%. These standard machine learning competitions show the potential for brain-based mental illness predictions, as they are able to evaluate data with accurate, unbiased predictive power.
7.4 | Benefits of leveraging dynamic connectivity features

Until recently, functional connectivity has been assumed to be relatively stable over the scanning time (usually several minutes). While convenient for analysis and interpretation purposes, this oversimplified assumption was recently challenged by several studies focused on time-varying multivariate connectivity patterns (Sakoğlu et al., 2010), as well as others focusing on time-frequency analysis methods (C. Chang & Glover, 2010). Several other studies have also delved into the time-resolved connectivity measures and their successful applications in identifying biomarkers using dynamic connectivity features (Allen et al., 2014; Calhoun, Miller, Pearson, & Adali, 2014; Du et al., 2016; Rashid et al., 2018; Rashid, Damaraju, Pearson, & Calhoun, 2014; Zalesky & Breakspear, 2015). These studies reported that brain functional connectivity can vary within a short period (e.g., tens of seconds), and can successfully capture the connectivity disruptions in a disease population.

Only a few studies have utilized dynamic brain connectivity features to predict mental illness. Using static and dynamic connectivity features, Rashid and colleagues developed a classification framework to predict SZ, bipolar, and healthy subjects (Rashid et al., 2016). The classification performance measures among static, dynamic and combined static and dynamic connectivity features were compared using a 10-fold CV framework. The results showed that dynamic FNC (classification accuracy: 84.28%) significantly outperformed the static FNC (classification accuracy: 59.12%), suggesting that dynamic patterns in functional connectivity might provide distinct and more information over the static FNC. Another study used a novel recurrent neural network (RNN) approach (i.e., a family of deep learning) to measure temporal dynamics and dependencies among brain networks (X.-H. Wang, Jiao, & Li, 2018). Using SVM classifier and LOO/10-fold CV, another study classified ADHD children using data from the ADHD-200 Consortium. Specifically, the temporal variability between intrinsic connectivity networks and the demographic and covariate variables were utilized as features. The LOOCV achieved total accuracy of 78.75, while the 10-folds CVs achieved average prediction accuracy of 75.54% ± 1.34, showing evidence of accurate prediction of ADHD using temporal dynamics and SVM classifier. Further, Demirtas and colleagues used SVM classifier and global spatiotemporal measures, static FNC and variability in FC, and reported that the combined static FC and variability in FC provided a classification accuracy of 81% (sensitivity: 81%; specificity: 81%) for classification of MDD (Demirtaş et al., 2016).

7.5 | Fusion of dynamic connectivity and other data types

Dynamic FNC measures estimated from fMRI data can be further integrated with other data types and modalities, for example, genomic and structural MRI data, to leverage inter-modality based features for disease characterization. In a novel imaging-genomic framework, Rashid and colleagues have recently modeled the association between dynamic FNC states and genomic features to examine the SZ-related inter-modality abnormalities (Rashid et al., 2019; Figure 17). Specifically, the parallel ICA algorithm (J. Liu et al., 2009) was utilized to combine genetic variants (i.e., single nucleotide polymorphism [SNP]) and functional features from fMRI data as subject-specific states that are revealed from the dynamic FNC data using a sliding window and clustering approach (Allen et al., 2014; Rashid et al., 2014) to distinguish SZ from healthy individuals. Results identified a significant association between a SNP component (defined by large clusters of functionally related SNPs statistically correlated with phenotype components) and dynamic FNC component (defined by clusters of related connectivity links among distant brain regions distributed across discrete dynamic states, and statistically correlated with genomic components) in SZ. Moreover, the polygenetic risk score (PRS) for SZ (computed as a linearly weighted sum of the genotype profiles with weights derived from the odds ratios of the psychiatric genomics consortium [PGC]) showed negative correlation with the significant dynamic FNC component, which were mostly present within a state that exhibited a lower occupancy rate in individuals with SZ compared with HC, therefore identifying a potential dynamic FNC imaging biomarker for SZ. Another recent study by Abrol and colleagues proposed an mCCA+jICA framework to fuse dynamic FNC from fMRI data and gray matter maps from structural MRI data (Abrol, Rashid, Rachakonda, Damaraju, & Calhoun, 2017). The framework identified the associated changes in both modalities, highlighting significantly disrupted links between dynamic FNC and gray matter volumes in SZ patients. Results from this study reported significant group differences in gray matter maps, particularly in the superior parietal lobule, precuneus, postcentral gyrus, medial/superior frontal gyrus, superior/middle temporal gyrus, insula and fusiform gyrus. Further, results also highlighted alterations in several inter-regional connectivity strength in SZ patients. In the field of brain-based prediction of mental illness, fusion approaches using dFNC and other features could increase the discriminative power of the models, although future studies are required to confirm their utilities in this regard.

8 | SUMMARY AND CONCLUSION

Recent brain-based mental illness predictome studies have shown promising results, although some results require further validation due to small sample sizes. While many advanced algorithms have been developed and applied in the field of mental illness prediction, there exists many challenging issues which must be further resolved prior to their applications in clinical settings. In this work, we comprehensively review existing brain-based prediction studies in several mental illnesses such as SZ, depressive disorders (i.e., MDD and BP), ASD, ADHD, SAD, OCD, PTSD, and substance disorder. We also highlight a number of existing approaches and future research directions. A major challenge in the field is the prediction of phenotypic heterogeneity that characterizes psychiatric disorders. However, recent approaches
have started to address the disease subtypes. This can improve disease prediction, provide biological support for existing categories or support the revision of existing diagnostic categories. Another major challenge is the relatively small sample size reported across most studies. Without more robust validation, it is unclear how generalizable these results will be when applied to an independent dataset. However, recent data-sharing initiatives have started to improve the sample size issue by offering adequate data to develop more robust and improved prediction models.

While brain-based classification has proven challenging, there has been considerable progress made in recent years. With the accelerating growth of large volumes of patient data and data-sharing initiatives in the field of neuroimaging and medicine, we anticipate diagnostic tools operating on comprehensive biomarker profile accessed from multiple modalities will be available for specific use cases in the near future. With more sophisticated deep learning models integrated with large-scale data, we believe that predictive modeling tools will soon transition from the "proof-of-concept" stage to the "ready for clinical implementation" stage. Further, while patient characteristics appear to be more homogeneous within relatively small samples (Schnack & Kahn, 2016), failing to capture disease-specific variability, with the power of "big" brain data and advanced machine learning algorithms, it is now possible to explore the heterogeneity within a disease (i.e., disease sub-types). It is a common practice for clinicians to consider diagnostic homogeneity (i.e., patients with similar clinical symptoms belong to the same broader diagnostic category). We can also evaluate this within imaging data using, for example, N-way clustering approaches to identify subgroups of homogeneous data followed by evaluation of clinical phenotypes. One recent example shows this in SZ and finds enhanced sensitivity to group variabilities, with the power of "big" brain data and advanced machine learning algorithms, it is now possible to explore the heterogeneity within a disease (i.e., disease sub-types). It is a common practice for clinicians to consider diagnostic homogeneity (i.e., patients with similar clinical symptoms belong to the same broader diagnostic category).

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