Multivariate pattern analysis strategies in detection of remitted major depressive disorder using resting state functional connectivity

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Abstract

Understanding abnormal resting-state functional connectivity of distributed brain networks may aid in probing and targeting mechanisms involved in major depressive disorder (MDD). To date, few studies have used resting state functional magnetic resonance imaging (rs-fMRI) to attempt to discriminate individuals with MDD from individuals without MDD, and to our knowledge no investigations have examined a remitted (r) population. In this study, we examined the efficiency of support vector machine (SVM) classifier to successfully discriminate rMDD individuals from healthy controls (HCs) in a narrow early-adult age range. We empirically evaluated four feature selection methods including multivariate Least Absolute Shrinkage and Selection Operator (LASSO) and Elastic Net feature selection algorithms. Our results showed that SVM classification with Elastic Net feature selection achieved the highest classification accuracy of 76.1% (sensitivity of 81.5% and specificity of 68.9%) by leave-one-out cross-validation across subjects from a dataset consisting of 38 rMDD individuals and 29 healthy controls. The highest discriminating functional connections were between the left amygdala, left posterior cingulate cortex, bilateral dorso-lateral prefrontal cortex, and right ventral striatum. These appear to be key nodes in the etiopathophysiology of MDD, within and between default mode, salience and cognitive control networks. This technique demonstrates early promise for using rs-fMRI connectivity as a putative neurobiological marker capable of distinguishing between individuals with and without rMDD. These methods may be extended to periods of risk prior to illness onset, thereby allowing for earlier diagnosis, prevention, and intervention.

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1. Introduction

Major depressive disorder (MDD) is a heterogeneous disorder characterized by variable patterns of nine key symptoms, often in episodic patterns across an individual’s lifetime, parallel to the waxing and waning observed in chronic illnesses such as multiple sclerosis (APA, 1994) with relapsing–remitting and relapsing–progressive patterns. Despite awareness that symptom profiles and illness course patterns vary widely across (and perhaps even within) individuals with MDD, prior studies have typically used the full diagnostic spectrum (studies of those with any and all patterns of MDD) and the broadest range in age and experience of illness in cross sectional studies. These prior, broad-stroke, heterogeneous studies may have led to increased type I error with relatively small samples and publication bias or diffusion of important, specific effects, increasing type II error. Together, these would dilute both inferential capability and replication. More recently, subtypes of MDD have been pursued with the emergence of some larger studies including more clinically and demographically homogeneous samples (Korgaonkar et al., 2014). To this end, the present study has strived to constrain a number of features with known impact on brain function in MDD varying from small (e.g., medications, subtypes of MDD) to medium (active symptoms) to very large (age, development) effect sizes. As such, we studied early course MDD in the remitted state among a sample of late-adolescents who were medication-free at the time of scan in order to reduce sources of heterogeneity for between group comparisons. These methodological controls increased homogeneity and have resulted in an emerging model for how mood disorders might have distinct trait/risk features, scar patterns, symptom perturbations, and chronic burden/scar components (Weisenbach et al., 2014; Peters et al., 2016; Votruba and Langenecker, 2013).

Studying individuals with rMDD enables a unique examination of potential trait-based mechanisms of depression and depression relapse (e.g. Marchetti et al., 2012). One method for understanding trait-based markers for MDD involves studying network function through measurements of network connectivity. Disrupted network connectivity...
has been documented among individuals within a major depressive episode (Greicius et al., 2007; Connolly et al., 2013), including within and between key nodes of posterior cingulate cortex (PCC), medial prefrontal cortex (mPFC), inferior parietal cortex (IPC; Hamilton et al., 2012), amygdala (Siegle et al., 2007), anterior insula, dorsal ACC (Strigo et al., 2008; Bricelj et al., 2013), pregenual ACC (Horn et al., 2010), DLPFC (Bench et al., 1992; Mayberg et al., 1999; Jacobs et al., 2014), and hippocampus (Cao et al., 2012; Sambataro et al., 2013).

Although alterations in brain functional connectivity have been demonstrated in MDD, most of these studies have focused on group level analysis. However, there is substantial interest in identifying single subject biomarkers that are clinically applicable as diagnostic or prognostic tools (Mossner et al., 2007; Atluri et al., 2013; Schneider and Pervolovic, 2013). Recent studies (Klöppel et al., 2011; Mourao-Miranda et al., 2012; Orru et al., 2012; Zarogiani et al., 2013; Haller et al., 2014; Sundermann et al., 2014) have begun to investigate diagnostic classification of mental disorders using rs-fMRI and multivariate pattern analysis (MVPA). Particularly, pioneering work has examined the clinical applicability of examining resting state data using MVPA (Cradock et al., 2009). MVPA algorithms provide a framework for disease state prediction in which the final goal is to predict the presence or absence of a disease based on observed functional connections.

The patterns are learned from multivariate data given predetermined categories, and performance is measured by the prediction accuracy obtained when classifying a new case. Support Vector Machine (SVM) has been used as a classification model often in fMRI research that may offer better prediction accuracy and is less sensitive to noise than alternative MVPA approaches (Mitchell et al., 2004; Chen et al., 2006; LaConte et al., 2007; Mourao-Miranda et al., 2005). In practice, building a robust, generalizable classification model can be challenging because the number of features far exceeds the number of data observations. To avoid model overfitting, there is a need to select informative features before building a classification model. In the context of rs-fMRI research, the features are the functional connections between two regions (voxels).

The objective of feature selection include the reduction of prediction error and the improved interpretability of a MVPA model (Guyon and Elisseeff, 2003a; De Martino et al., 2008; Mourao-Miranda et al., 2005). Several filter and wrapper approaches (Cradock et al., 2009; Zeng et al., 2014; Cao et al., 2014) have been used based on univariate t-test, probability density function and recursive feature elimination (Guyon and Elisseeff, 2003b) in fMRI research. Embedded methods learn which features best contribute to the accuracy of the model while the model is being created. The most common type of embedded feature selection methods are regularization methods. In recent years, regularized embedded methods such as LASSO and Elastic Net have demonstrated good effectiveness and sensitivity (Mwangi et al., 2013) in neuroimaging machine learning tasks such as Alzheimer’s disease (AD) classification (Casanova et al., 2011; Rao et al., 2011; Shen et al., 2010), treatment response predictions in Attention Deficit Hyperactivity Disorder (Marquand et al., 2012) and Autism Spectrum Disorder classification (Duchesnay et al., 2011). In the context of rs-fMRI, a few studies have applied SVM (Cradock et al., 2009; Cao et al., 2014) in identifying patients with MDD. With task-based data, machine learning using a Gaussian Process Classifier has been used to discriminate adolescents at high risk for mood disorders from healthy adolescents (Mourao-Miranda et al., 2012). However, to our knowledge, rs-fMRI connectivity data have not been explored using machine learning among patients in the remitted state of MDD.

In this research, we built a sparse MVPA framework combining regularized Elastic Net feature selection algorithm and a linear SVM. The advantage of Elastic Net regularization penalty over filter approaches is that it conducts automatic variable selection and continuous shrinkage simultaneously, and selects a group of correlated variables. This feature selection strategy is a state-of-the-art representative of recent advances in L1/L2-constraint based methods. We compared Elastic Net with two filter approaches t-test and Wilcoxon rank sum and performed these evaluations using leave-one-out cross validation in the context of a study of resting state functional connectivity in remitted major depressive disorder (rMDD). This design allowed us to examine network differences in the absence of the state effects of active illness. We hypothesized that building a MVPA framework by employing feature selection strategies combined with SVM would be successful in identifying the discriminant functional connections that predicted prior history of rMDD.

2. Methods

2.1. Participants

Participants were recruited from the University of Michigan (UM) and the University of Illinois at Chicago (UIC) using flyers and multiple forms of posting on the internet. All participants completed an identical assessment protocol, including the Diagnostic Interview for Genetic Studies (DIGS; Nurnberger et al., 1994), the Hamilton Depression Scale (Ham-D; 34 Hamilton, 1960), and a targeted neuropsychological and fMRI battery (not reported here). Participants were considered remitted from MDD if they previously met criteria for at least one major depressive episode (MDE), did not meet current criteria for an MDE in the last three months (Mean 2.5 years well), and currently scored below a 7 on the Ham-D (administered during the phone screen and during the initial diagnostic interview). HCs could not meet current or past criteria (Never Mentally II, NMI) for MDD or any other Axis I or II psychiatric disorder and had no first degree relatives with a history of psychiatric illness. In addition, participants were required to be medication free for a period of 30 days prior to the scan and those with substance abuse or dependence within the past six months were excluded.

Diagnosis of past MDD or NMI was confirmed using a modified Family Interview for Genetic Studies completed with a parent, guardian, or older sibling (Nurnberger et al., 1994). The final sample included 38 rMDD (17 UM, 21 UIC) and 29 NMI (16 UMC, 13 UIC) between the ages of 18–23 years (66% Female). None of the remitted (r)MDD were taking medications at the time of scan or for the past 30 days (and 21 had never taken any psychotropic medication, 33 had a history of psychotherapy). The rMDD (aged 18–23, modal depressive episodes = 1, modal years well = 4) were compared with data from 29 NMI. Participant demographics and clinical characteristics are presented in Table 1.

| Site | N | Site | N |
|------|---|------|---|
| UM   | 38| UIC  | 29|
| UIC  | 17| UM   | 16|
| Sex  | 29| Age  | 20.97 (SD = 1.55)|
| F/M  | 6/13| M   | 14.34 (SD = 1.21)|
| Years Education | 13 Yes/25 No |
| 14 | 35 | 14 | 90 |
| 1.92 (SD = 1.25) | 1.53 (SD = 1.40) |
| 1.55 (SD = 1.21) | 1.34 (SD = 1.40) |
| 1.25 (SD = 1.21) | 1.25 (SD = 1.40) |
| NA | NA |
| NA | 4.91 |

rs-fMRI data from two 3.0 Tesla GE scanners were collected using eight bilateral seeds in the default mode network (DMN), salience network (SN) and cognitive control network (CCN). Table 2 provides the

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* Indicates that there are significant differences in HAM-D. No participants had any current medication use for at least the past 30 days.
Montreal Neurological Institute (MNI) coordinates of the ROI seeds. Seeds were derived based on previous literature examining resting state connectivity of the amygdala (Fox et al., 2009; Pannekoek et al., 2013), PCC (McCabe and Mishor, 2011; Bluhm et al., 2011), sgACC (Alexopoulos et al., 2012; Kelly et al., 2009), and anterior superior insula (Margulies et al., 2007; Siegle et al., 2006) in depression. All seeds were verified visually on an average anatomy of the first 55 subjects that participated in the study, and used 19 contiguous voxels (radius 2.9 mm).

### 2.3. rs-fMRI preprocessing

Data preprocessing occurred as follows: Slice timing was completed with SPM8 (http://www.fil.ion.ucl.ac.uk/spm/doc/) and motion detection algorithms were applied using FSL (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/). Coregistration of structural images to functional images was performed by regressing out white matter and cerebral spinal fluid signals (Tabachnick and Fidell, 2007). Motion parameters were regressed out (Behzadi et al., 2007). Based upon the recent literature (Behzadi et al., 2007; Jo et al., 2013), motion volumes were identified based on any TR to TR movement exceeding 1.5 mm and did not differ between groups. Those with significant movement were not included in this final reported sample. Global signal was not regressed due to collinearity violations with gray matter signal, problematic mis-estimates of anticorrelations (Power et al., 2012), and because it does not affect distance-micromovement relationships (Behzadi et al., 2007). Finally time-series were band-pass filtered over 0.01–0.10 Hz. Regions of Interest (ROIs; 2.9 mm radius) were defined in MNI space and spatially averaged time course data were extracted from these regions for each participant. Correlation coefficients were calculated between seed regions (Table 2) and transformed to z scores using a Fisher transformation.

### 2.4. Cross-correlation analysis

The rs-fMRI time series was detrended and mean centered. Physiologic correction was performed by regressing out white matter and cerebral spinal fluid signals (Tabachnick and Fidell, 2007). Motion parameters were regressed out (Behzadi et al., 2007). Based upon the recent literature (Behzadi et al., 2007; Jo et al., 2013), motion volumes were identified based on any TR to TR movement exceeding 1.5 mm and did not differ between groups. Those with significant movement were not included in this final reported sample. Global signal was not regressed due to collinearity violations with gray matter signal, problematic mis-estimates of anticorrelations (Power et al., 2012), and because it does not affect distance-micromovement relationships (Behzadi et al., 2007). Finally time-series were band-pass filtered over 0.01–0.10 Hz. Regions of Interest (ROIs; 2.9 mm radius) were defined in MNI space and spatially averaged time course data were extracted from these regions for each participant. Correlation coefficients were calculated between seed regions (Table 2) and transformed to z scores using a Fisher transformation.

### 2.5. Feature extraction

The rs-fMRI network was captured by a 16 × 16 symmetric matrix of nodes. We extracted the upper triangle elements of the functional connectivity matrix as classification features, i.e. the feature space for classification was spanned by the (16 × 15) / 2 = 120 dimensional feature vectors.

### 2.6. Classification algorithm

SVM classification (Vapnik, 1995) is a widely used method for binary classification in fMRI studies. SVMs are supervised learners that work in two steps. In the training step, a subset of the available data points as well as their associated classes is used to iteratively find a linear boundary or hyperplane that separates the two classes optimally. In the testing step, new, previously unobserved data points in the same space as the training points are classified depending on their position relative to the boundary (often the “case left out” in leave one out, small n analyses). For two classes, the SVM algorithm attempts to find a linear decision boundary (separating hyper plane) using the decision function $D(u) = \langle w, u \rangle + w_0$, where $w$ defines the linear decision boundary, and is chosen to optimize the boundaries defined by $D = +1$ and $D = -1$ (known as the margin) between the two class distribution. The decision function that is learned by a SVM is a linear combination of feature values in a particular feature space.

There are differences between different types of SVM, notably in how the relationship between the feature space and the original features (functional connections, in our case) is determined; a given choice of kernel function determines an (implicit) feature space in which a decision takes place.

### 2.7. Feature selection algorithms

In neuroimaging studies, the number of features are often more than the number of observations (often less than 100), which causes curse-of-dimensionality and small-n-large-p effects. The unimportant features may result in an over fitting problem in machine learning, and therefore reduces model prediction accuracy and generalization ability. We considered two embedded-methods: LASSO and the Elastic Net (Tibshirani, 1996; Zou and Hastie, 2005). The popular LASSO regression (Tibshirani, 1996) method minimizes the Residual Sum of Squares (RSS), similar to Ordinary Least Squares (OLS) regression, but poses a constraint to the sum of the absolute values of the coefficients being less than a constant. This additional constraint is similar to that introduced in Ridge regression, where the constraint is to the sum of the squared values of the coefficients. This simple modification allows LASSO to perform also variable selection because the shrinkage of the coefficients is such that some coefficients can be shrunk exactly to zero. The LASSO computes model coefficient $\beta$ by minimizing the following function $R(\beta) + \lambda \sum_{j=1}^{p} |\beta_j|$, where $R(\beta)$ is the mean square error on the training set and $\|\beta\|_0 = \sum_{j=1}^{p} |\beta_j|$. $\lambda$ controls the degree of sparsity of the solution, i.e. the number of features selected.

Elastic Net (Zou and Hastie, 2005) is similar to LASSO. It differs in that the $l_1$ norm of $\beta$ is replaced by a combination of $l_1$ and $l_2$ norms. In this case, we minimize $R(\beta) + \lambda \text{Pen}(\beta)$, where $\text{Pen}(\beta) = \frac{1}{2} \sum_{j=1}^{p} \beta_j^2 + \alpha \sum_{j=1}^{p} |\beta_j|$, for $\alpha$ strictly between 0 and 1, and a nonnegative $\lambda$. The $\lambda$ parameter can be tuned in order to set the shrinkage level, and the higher the $\lambda$ is, and the more coefficients are shrunk to 0. Elastic Net is the same as LASSO when $\alpha = 1$. As $\alpha$ shrinks toward 0, Elastic Net approaches ridge regression. For other values of $\alpha$, the penalty term $P_1(\beta)$ interpolates between the $L_1$ norm of $\beta$ and the squared $L_2$ norm of $\beta$. The advantage of Elastic Net over LASSO is that the Elastic Net penalty completes automatic variable selection and continuous shrinkage simultaneously, and it can select from a group of correlated variables. It is especially useful for large $p$ small $n$ problems where the grouped variables situation is a particularly important concern (Hastie et al., 2000, 2001).

### 2.8. Feature selection and classification method

Due to our limited number of samples, we used a leave-one-out cross-validation (LOOCV) strategy to estimate the generalization ability of our classifier. We used SVM classifier with three strategies: 1) No
feature selection, which acted as a baseline to focus the specific contributions of feature selection. 2) Multivariate LASSO and Elastic Net based feature subset ranking. 3) Univariate t-test and Wilcoxon feature subset ranking.

Feature selection algorithms were implemented in MATLAB 2012b and were done in each cross validation fold. For Elastic Net, we have evaluated classification performances for different values of alpha, ranging from .1 to 1.0. When alpha = 1, it is Lasso and when alpha is between 0 and 1, it is Elastic net. Inside each LOOCV fold, 10-fold CV was used to select the best elastic net regularization parameter lambda ($\lambda$).

We also evaluated two filter-based algorithms such as Student’s t-test and Wilcoxon sum-rank test to rank the features which evaluate if each feature is significantly different between the two classes; these ranking algorithms were employed in each cross validation fold. We used the MATLAB Bioinformatics toolbox to compute these scoring functions.

Inside the LOOCV ($n = 67$), the features were selected based on a feature selection strategy and used as the final feature set for SVM classification. We used default parameter setting in Matlab SVM function for kernel and optimization method. For the selection of soft margin parameter C, we performed another 10-fold CV strategy for different kernel and optimization method. For the selection of soft margin parameter value C which produces highest accuracy in 10-fold CV and the feature (functional connections) contributions to classification. We computed the percentages of occurrences of features that contributed to identification of depressed patients across all iterations of the cross validation. The functional connectivities which appeared in the leave-one-out process more than half of the times were shown in Fig. 5, and indicated the most discriminative features between those with rMDD and HC.

We also performed a hold-out validation test by randomly selected nine samples (5 rMDD patients, 4 controls) as testing data set and remaining 58 samples as the training dataset. We obtained the best alpha and k parameters by re-running the LOOCV using only the training 58 subjects. Once we found the connectivities using 58 subjects with best alpha and k parameters, we used these connections as our final model and tested with the hold-out 9 subjects for prediction. The results are shown in Table 4.

2.9. Permutation test

The performance of the SVM classifier was evaluated using accuracy, sensitivity and specificity measures. To determine whether classification accuracy exceeded chance levels (50%), we performed permutation testing and derived a $p$-value. We permuted the class labels 1000 times (each time randomly assigning rMDD and HC labels to each pattern of functional connectivity values) and repeated the entire feature selection algorithm. We then counted the number of times the permuted test accuracy was higher than the one obtained for the true labels. Finally, we divided this number by 1000 and obtained a $p$-value for classification accuracies. For the permutation test the 10-fold CV and LOOCV structures were maintained.

3. Results

First we employed a two sample t-test using our data set to find any significant functional connectivity differences between healthy controls and rMDD. There were eight connections found to be significantly different among these two groups (Fig. 2). At a more conservative test, we also performed a FDR corrected t-test ($q = .2$), which found four significant group differences in connections (Fig. 2, left four panels).

It should be noted that univariate filter methods ranked all variables in terms of relevance, as measured by a score. In each CV fold, we selected the top $k$ FCs according to the score and reported the average accuracy over all CV folds. Fig. 3 illustrates the performance of t-test and Wilcoxon methods for different values of $k$. As the chart reveals, the highest accuracy was obtained when $k = 4$ for t-test and $k = 3$ for
Wilcoxon test. With higher $k$ after that, accuracy declined. In contrast, LASSO and Elastic Net regressions derived only important FCs, with a different number of important FCs in each CV fold. As such, the SVM classifier was trained with only the important predictors found in each CV. The average accuracy for different values of $\alpha$ parameters is reported. As illustrated in Fig. 4, the best accuracy (76.1%) was obtained when $\alpha = 0.5$. At $\alpha = 1$, Elastic Net becomes LASSO, and very few variables (mostly two features in each CV fold), and the accuracy is 67.1%.

Table 3 summarizes the performance of t-test, Wilcoxon, LASSO and Elastic Net using SVM classifier across 67 folds. Without any feature selection algorithm, the SVM classifier using LOOCV obtained overall accuracy of 45.1% (57.9% for patients and 28.3% for controls). In contrast, the embedded methods LASSO and Elastic Net algorithms discard the unimportant features by forcing them to zero, so there is no ranking of features. The highest classification accuracy of 76.1% was obtained by choosing the Elastic Net feature selection algorithm (81.5% for patients and 68.9% for healthy controls). t-Test and Wilcoxon methods also performed well, both with 71.6% accuracy (76.3% for patients and 65.5% for healthy controls).

The classification accuracies from holdout validation test are shown in Table 4. We found the consensus connectivities using 58 subjects with best alpha and $k$ parameters by carrying out a LOOCV. We used these connections as our final model and tested with the remaining 9 subjects for prediction. Our results showed that the both filter-based t-test and Elastic Net obtained accuracy of 77.8%.

The percentages of functional connections in all folds that contributed to identification of rMDD patients are shown in Fig. 5. We showed only those functional connections for the final model which appeared more than 50% inside LOOCV using 58 subjects. The most discriminative connections based on consensus functional connections were between the left PCC and right DLPFC and between the left amygdala and right VSs, among all feature selection algorithms. These connections were selected in all cross validation folds. The connections between left PCC and left DLPFC were selected more than 95% of the time using Elastic Net, Wilcoxon and t-test ranking algorithms. The connections between left amygdala and left VSs were observed in Elastic Net and t-test algorithms in more than 92% of cross validation folds. In addition, Elastic Net identified the connection between right PCC and left DLPFC in 61% of cross validation folds.

4. Discussion

The present study aimed to classify participants into the clinical psychiatric diagnostic categories for rMDD or NMI using SVM of rs-fMRI data. Sixteen nodes from the DMN, SN and CCN were chosen a priori based on previous research indicating their importance in MDD and rMDD (Jacobs et al., 2014; Tremblay et al., 2005; Kaiser et al., 2015) and theories show that these three networks comprise the core neurocognitive networks (Raichle et al., 2001; Seeley et al., 2007; Menon, 2011). We then applied SVM, using four different feature selection algorithms, to identify the top ranked features (connections between two nodes) that distinguished the rMDD from the HC group. In support of our hypothesis, we were able to use SVM of rs-fMRI data to discriminate these participants, even in the remitted state. We found that the most discriminative connection was between the left PCC and the right DLPFC, followed by the left amygdala and the right superior...
ventral striatum connection. These two connections were selected in 100% of training folds for all feature selection methods.

Our results provide new support in discrimination of rMDD, extending beyond previous studies that have been able to use machine learning classification, including SVM, to discriminate active MDD (aMDD) from HC using rs-fMRI data (Guo et al., 2012; Guo et al., 2014). For example, using SVM with leave-one-out cross validation applied to whole-brain rsfMRI data, active MDD were distinguished from HC with about 84% classification accuracy (Cao et al., 2014). A similar result (93% accuracy) was obtained in predicting active MDD using 15 pre-determined regions of interest (Craddock et al., 2009). SVM classification performance here (76.1% accuracy) was similar, but not as high as reported in the above mentioned studies. However, it is important to note that we did not employ separation of active MDD from NMI. The results of prior studies of aMDD could capture features related to active illness state, trait features, and potential scar/repetitive scar features, strengthening results. Here, our results could only be related to trait and early scar features, weakening the number of predictors, but strengthening the meaning and actual practical use. In addition, our feature selection algorithms are different and we used a larger sample which can result in more conservative estimates. In the future, we intend to explore other feature selection strategies to use strategies like recursive feature elimination and sparse logistic regression. It is worthwhile to note that, in a clinical setting, when our goal is to find the pattern of functional connectivity that accurately predicts whether a subject suffers from rMDD, we expect a classification performance to be 95–100% accuracy (Orru et al., 2012). This is a promising initial step that can be followed up using additional predictors and moving to at-risk samples.

### 4.1. DMN–CCN connectivity

One of our main discriminant connections for rMDD was between the left PCC and right DLPFC. These regions are part of the DMN and the CCN, respectively, and reflect meta-analytic findings of hyperconnectivity/hyperactivity of posterior default network components including the PCC with lateral prefrontal areas (Sundermann et al., 2014; Kaiser et al., 2015). Our results also lend support to previous research that found disturbances in the PCC and DMN, also in people with active MDD (Hamilton et al., 2012; Pizzagalli, 2011), including increased connectivity in the PCC in patients with active MDD (Sambataro et al., 2013). As we found these patterns for the first time in young adults with rMDD, it lends credence to the argument that this dysfunctional connection is not related to active symptoms, and it is present early in the course of the disease. A dysfunctional DLPFC may represent a precursor to the hyperactive midline activity reported in MDD, and connections to the DMN may represent a vulnerability marker in people at-risk for MDD (Marchetti et al., 2012). It is possible that cognitive risk factors, including rumination and poor attentional control, are a direct byproduct of DMN hyperconnectivity, and are associated with maladaptive self-focus. Consistent with this argument, individuals with MDD have a tendency to attend to internal at the expense of external stimuli (Greicius et al., 2007; Hamilton et al., 2011; Broyd et al., 2009), and aberrant connectivity within the DMN has been associated with increased rumination and brooding in depression (Hamilton et al., 2011; Berman et al., 2010; Jacobs et al., 2014; Zhu et al., 2012) and decreased sustained attention (Jacobs et al., 2014).

### 4.2. SN connectivity

Our second discriminant connection for rMDD was between the left amygdala and right ventral striatum. Within the three core network model, both the ventral striatum and the amygdala are considered part of the salience (and emotion) network. Connections between limbic structures such as the amygdala and the VS highlight how and where affective processes could influence action (Mogenson et al., 1980; Mogenson and Nielsen, 1984). The ventral striatum includes the nucleus accumbens, which is a critical part of the mesolimbic dopaminergic reward system. Dopamine function in reward-related processes, and in particular disruption of these in animal models, led to the development of the anhedonia hypothesis by Wise and colleagues (Wise, 1982, 1985). Subsequent observations of VS dysfunction patients with MDD are interpreted in light of this hypothesis (Epstein et al., 2006; Keedwell et al., 2005; Pizzagalli et al., 2009), including emergence early in the course of the illness.

In contrast, the amygdala is important for the detection of salient stimuli in the environment, including facial expressions, often of a negative or threatening valence (Adolphs et al., 1994; Fu et al., 2008). It functions as a low signal detection system to orient the organism toward such potentially important stimuli (Holland and Gallagher, 1999; Whalen, 1998) so that the correct behavior can be enacted. Amygdala activity has been shown to be heightened in active MDD (Fu et al., 2008; Shelton et al., 2010; Surguladze et al., 2005). SN dysfunction and disrupted connectivity could thus contribute to emotional regulation difficulties in MDD via biasing attention to emotional stimuli, even those below the level of conscious awareness (Victor et al.,

### Table 3

SVM Classification accuracies in discriminating between rMDD patients and controls using LOOCV.

| Methods      | Accuracy (%) | Sensitivity (%) | Specificity (%) | Permutation test* | p-Value |
|--------------|--------------|----------------|-----------------|-------------------|---------|
| All features | 44.4         | 60.0           | 25.0            | >.05              |         |
| Elastic Net  | 77.8         | 80.0           | 75.0            | <.05              |         |
| t-Test       | 66.7         | 80.0           | 50.0            | >.05              |         |

* Permutation test indicates whether the accuracy exceeds chance levels (50%).
2010), and via connections to autonomic regions involved in the psychophysiological emotional response (Drevets et al., 2008; Price and Drevets, 2010).

Given previous findings indicating the importance of sgACC connectivity in active MDD as a predictor of depression refractoriness and severity (Greicius et al., 2007; Connolly et al., 2013; Mayberg et al., 1999), it may be surprising that the current study did not identify sgACC connectivity as a top feature. It is possible that sgACC activity is a state effect of MDD, as sgACC hyperactivity has been reported to normalize following treatment of MDD, and is even present in sad mood induction (Fu et al., 2008; Mayberg et al., 1999; Sheline et al., 2010; Victor et al., 2013). In terms of connectivity, our results failed to support previous findings of studies using machine learning that the connectivity of the sgACC is able to differentiate active MDD patients from HC s with a high degree of accuracy (e.g. Zeng et al., 2014). However our results do support findings that the amygdala is hyperconnected in MDD (e.g. Jin et al., 2011), and suggest that this may be either a risk factor or trait effect, as it persists into the remitted state.

4.3. Lack of SN–CCN connectivity predictor

Studies that have investigated abnormalities in regional functional connectivity in active MDD have reported reduced connectivity between cortical regulatory areas and the amygdala during task performance (Carballedo et al., 2011; Dannowski et al., 2009; Siegel et al., 2007; Moses-Kolko et al., 2010), and activation in the nucleus accumbens as well as the amygdala during successful inhibition on a cognitive control task has been found to predict post-treatment improvement in depressive symptoms (Langenecker et al., 2007). Anomalous functional connections may be reflective of disrupted interactions between the SN and CCN (Dannowski et al., 2009). The DLPCF and VLPCF/insula are important for both explicit and implicit cognitive control of limbic regions and the correlated emotional responses (Langenecker et al., 2014). As such, we were surprised that there was not a significant discriminant feature connection between a CCN region and a SN region, such as the amygdala. Thus, anomalous functional connectivity between regions of the salience and cognitive control networks in MDD may be a state effect, or may emerge with longer standing illness course (more episodes).

4.4. Limitations

Multivariate pattern classification of resting state functional MRI data is a challenging task due to small samples with expensive data collection, noisy and high dimensionality of the data, and individual variability. We note several limitations in the current study. The first is a lack of an evaluation data set with which to test our methods and confirm the findings. Future research can confirm classification results with larger sample sizes and/or multicenter imaging data. The second limitation is that we used seed-based functional connectivity patterns. Using a whole brain functional connectivity measure may improve our classification accuracy. There is need for acquiring additional neuroimaging evidence of brain abnormalities including differences in structure that may discriminate individuals with MDD (active or remitted) from HCs. Using the current method, we cannot infer the directionality of connections, and therefore future research could use measures of effective connectivity such as Granger causality (Van den Heuvel and Hulshoff Pol, 2010). Another limitation is that the present study was cross-sectional, therefore we cannot dissociate the scar effects of illness from the risk factors for MDD. We will be following this sample longitudinally to determine which participants relapse, and which features of brain connectivity during remission can be used to classify individuals into relapsing or resilient groups, similar to what has been attempted using structural MRI to predict treatment response in MDD using SVM (Gong et al., 2011). Knowing which individuals are more likely to relapse will allow for more intensive therapies to be targeted to this group in order to avoid the chronic scarring effects of illness. It should be noted that while all of our sample were medication free for the previous month, only some of our sample were medication naïve, so we were unable to dissociate the effects of previous medication use. Future research can combine resting state functional connectivity and structural abnormalities to obtain more reliable clinical diagnosis of MDD. Finally, it is unfortunate that none of the connections observed with the SVM approach were unique from those captured in traditional univariate analyses.

5. Conclusions and future directions

The present study is the first to demonstrate that new SVM techniques can be used with rs-fMRI data to distinguish individuals with a history of MDD from controls with reasonably good accuracy. Efficient information and emotion processing requires coordination between a series of networks, even in the resting state (Di and Biswal, 2013; Spreng et al., 2013). Impaired internetwork connectivity at rest, even for individuals in the remitted state of MDD, were strong predictors of illness history. Our data suggests that there is remarkable potential in combining rs-fMRI data with machine learning-based techniques for advancing our understanding of network-level differences in MDD. It is hoped that machine learning techniques such as those used in the present study will develop further to be able to identify biomarkers that can inform us about individual prognosis such as relapse and thus guide therapeutic decisions and assist in psychiatric research into MDD.

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