

UvA-DARE (Digital Academic Repository)

Meta-Analysis of MicroRNAs Dysregulated in the Hippocampal Dentate Gyrus of Animal Models of Epilepsy

Srivastava, P.K.; Roncon, P.; Lukasiuk, K.; Gorter, J.A.; Aronica, E.; Pitkänen, A.; Petretto, E.; Johnson, M.R.; Simonato, M. DOI 10.1523/ENEURO.0152-17.2017

Publication date 2017 **Document Version** Final published version Published in eNeuro License

CC

Link to publication

Citation for published version (APA):

Srivastava, P. K., Roncon, P., Lukasiuk, K., Gorter, J. A., Aronica, E., Pitkänen, A., Petretto, E., Johnson, M. R., & Simonato, M. (2017). Meta-Analysis of MicroRNAs Dysregulated in the Hippocampal Dentate Gyrus of Animal Models of Epilepsy. eNeuro, 4(6), [e0152]. https://doi.org/10.1523/ENEURO.0152-17.2017

General rights

It is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), other than for strictly personal, individual use, unless the work is under an open content license (like Creative Commons).

Disclaimer/Complaints regulations

If you believe that digital publication of certain material infringes any of your rights or (privacy) interests, please let the Library know, stating your reasons. In case of a legitimate complaint, the Library will make the material inaccessible and/or remove it from the website. Please Ask the Library: https://uba.uva.nl/en/contact, or a letter to: Library of the University of Amsterdam, Secretariat, Singel 425, 1012 WP Amsterdam, The Netherlands. You will be contacted as soon as possible. UvA-DARE is a service provided by the library of the University of Amsterdam (https://dare.uva.nl)

Meta-Analysis of MicroRNAs Dysregulated in the Hippocampal Dentate Gyrus of Animal Models of Epilepsy

Prashant K. Srivastava,^{1,*} Paolo Roncon,^{2,*} Katarzyna Lukasiuk,³ Jan A. Gorter,⁴ Eleonora Aronica,⁵ Asla Pitkänen,⁶ Enrico Petretto,⁷ Michael R. Johnson,^{1,†} and Michele Simonato^{2,8,9,10†}

DOI:http://dx.doi.org/10.1523/ENEURO.0152-17.2017

¹Division of Brain Sciences, Imperial College London, Charing Cross Hospital, W12 0NN London, United Kingdom, ²Division of Neuroscience, University Vita-Salute San Raffaele, Milan 20132, Italy, ³Nencki Institute of Experimental Biology, Polish Academy of Sciences, 02-093 Warsaw, Poland, ⁴Swammerdam Institute for Life Sciences, Center for Neuroscience University of Amsterdam, 1105 Amsterdam, The Netherlands, ⁵Department of (Neuro)Pathology, Academic Medical Center and SEIN – Stichting Epilepsie Instellingen Nederland, The Netherlands, ⁶Department of Neurobiology, A.I. Virtanen Institute for Molecular Sciences University of Eastern Finland, FIN-70 211 Kuopio, Finland, ⁷Duke-NUS Medical School, Singapore 169857, Singapore, ⁸Department of Medical Sciences, Section of Pharmacology, University of Ferrara, Ferrara, Italy, ⁹National Institute of Neuroscience, Italy, and ¹⁰Laboratory for Technologies of Advanced Therapies (LTTA), Ferrara 44121, Italy

Abstract

The identification of mechanisms transforming normal to seizure-generating tissue after brain injury is key to developing new antiepileptogenic treatments. MicroRNAs (miRNAs) may act as regulators and potential treatment targets for epileptogenesis. Here, we undertook a meta-analysis of changes in miRNA expression in the hippocampal dentate gyrus (DG) following an epileptogenic insult in three epilepsy models. We identified 26 miRNAs significantly differentially expressed during epileptogenesis, and five differentially expressed in chronic epilepsy. Of these, 13 were not identified in any of the individual studies. To assess the role of these miRNAs, we predicted their mRNA targets and then filtered the list to include only target genes expressed in DG and negatively correlated with miRNA expression. Functional enrichment analysis of mRNA targets of miRNAs dysregulated during epileptogenesis suggested a role for molecular processes related to inflammation and synaptic function. Our results identify new miRNAs associated with epileptogenesis from existing data, highlighting the utility of meta-analysis in maximizing value from preclinical data.

Key words: dentate gyrus; epilepsy; hippocampus; meta-analysis; miRNA; mRNA

Significance Statement

Meta-analyses of data from human research studies are an invaluable tool, and the methods to conduct these investigations are well established. However, meta-analyses of preclinical data are rarely undertaken, due to the typically small sample sizes and the substantial heterogeneity between studies. We implemented a meta-analysis of microRNA (miRNA) expression changes in animal studies of epilepsy. This is the first study of its kind in the field of epilepsy and one of the first in preclinical research. Our analyses identify new miRNAs associated with epileptogenesis and epilepsy, highlighting common mechanisms across different animal models. These miRNAs and their predicted effects on gene expression generate new hypotheses about the causes of epilepsy that will prompt new studies in the field.

Received April 28, 2017; accepted November 22, 2017; First published December 13, 2017.

The authors declare no competing financial interests.

Authors contributions: P.K.S., P.R., M.R.J., and M.S. performed research; E.P., K.L., J.A.G., E.A., and A.P. designed research; P.K.S., P.R., M.R.J., and M.S. wrote the paper.

Introduction

Epilepsy is a serious, common neurologic disorder primarily characterized by the occurrence of spontaneous seizures. The treatment of epilepsy remains as one of the major unmet medical needs in neurology because, despite of over 20 antiepileptic drugs on the market, seizures are not controlled in about one third of the patients. The most common form of epilepsy in adults originates in temporal structures of the brain (temporal lobe epilepsy, TLE; Hauser et al., 1993). Epilepsy (TLE in particular) frequently arises as a consequence of brain injury ("acquired epilepsy"). While acquired epilepsies are in principle preventable by the therapeutic targeting of molecular processes underpinning their development (i.e., antiepileptogenic therapies), there are currently no established treatment options for halting the transformation of normal brain tissue to epileptic (Simonato et al., 2014). Identification of the mechanisms underlying epileptogenesis would therefore facilitate the identification of therapies for preventing the development of epilepsy and may inform new strategies for overcoming drug resistance in epilepsy more generally (Simonato et al., 2012; 2013; 2014).

Recent studies have suggested that microRNAs (miRNAs) play an important role in the pathogenesis of acquired epilepsy and may represent novel therapeutic targets (Jimenez-Mateos et al., 2012; Brennan et al., 2016; for review, see Cattani et al., 2016; Henshall et al., 2016). miRNAs are a family of small (21-25 nucleotides) noncoding RNAs, which can modulate various cellular and biological processes by degrading or repressing translation of specific mRNAs (Bartel, 2004, Guo et al., 2010). In systems analysis, miRNAs and their gene targets are described as following a "many-to-many" data model, such that each miRNA may regulate many transcripts and a single transcript may be regulated by many miRNAs (Ebert and Sharp, 2012). miRNAs have been implicated in various neuronal functions that are relevant in the pathogenesis of neurologic diseases, including epilepsy (Tan et al., 2013; Rajman et al., 2017; for review, see Karnati et al., 2015).

This work was supported by funding from the European Union's Seventh Framework Programme (FP7/2007-2013) under Grant Agreement 602102 (EPITARGET; to M.R.J., M.S., E.P., K.L., E.A., A.P.), the Imperial College National Institute for Health Research Biomedical Research Centre Scheme (M.R.J., E.P.), and the Polish Ministry of Science and Education Grant W19/ 7.PR/2014 (to K.L.).

*P.K.S. and P.R. contributed equally to this work.

[†]M.R.J. and M.S. share the position of senior author and Authors who analyzed data are P.K.S., P.R., M.R.J., and M.S.

Acknowledgements: We thank Dr. Ray Dingledine (Emory University, Atlanta, GA) for helpful discussions and for critically reading this manuscript.

Correspondence should be addressed to either of the following: Prof. Michele Simonato, Department of Medical Sciences, Section of Pharmacology, University of Ferrara, Via Fossato di Mortara 17-19, 44121 Ferrara, Italy, E-mail: michele.simonato@unife.it; or Prof. Michael Johnson, Division of Brain Sciences, Imperial College London, Charing Cross Hospital, W12 0NN London, United Kingdom, E-mail: m.johnson@imperial.ac.uk.

DOI:http://dx.doi.org/10.1523/ENEURO.0152-17.2017

Copyright © 2017 Srivastava et al.

Interpretation of data investigating the dysregulation of miRNAs in the brains of patients with epilepsy is challenged by the absence of appropriate human control brain tissue (Roncon et al., 2016). Research on the role of miRNAs in epilepsy has therefore focused on the use of experimental models of epilepsy, revealing changes in the expression of hippocampal miRNAs at different stages of the epileptogenic process (Henshall et al., 2016). However, methodological differences between the various preclinical animal models of epilepsy have made comparisons between studies difficult and the identification of common pathways dysregulated in epileptogenesis and epilepsy problematic. The current preclinical miRNA studies vary in multiple parameters, including brain region analyzed, animal model, sample size, microarray platform and analysis technique. Moreover, these studies are generally substantially underpowered to reliably detect modest changes in miRNA expression.

One particular concern is that analysis of large brain regions (like the hippocampus or the cortex) across different studies may confound interpretation and comparison because of variable cellular composition (e.g., relating to variable neuronal loss, astrocytosis, microgliosis, etc.). One way to address this issue could be to focus on a specific cell population. In this respect, dentate gyrus (DG) granule cells (GCs) seem particularly attractive as a target of analysis as the DG GC layer is a compact layer of (almost) identical cells, facilitating the dissection of a nearly pure cell population (GCs). Moreover, the DG has been traditionally described as a "gate" to inhibit hippocampal overexcitation (Chevaleyre and Siegelbaum, 2010) and recently, this hypothesis found support from new technologies; optogenetic GC hyperpolarization was found to stop spontaneous seizures, whereas optogenetic GC activation exacerbated spontaneous seizures, and activating GCs in nonepileptic animals evoked acute seizures (Krook-Magnuson et al., 2015). Finally, the DG is known to undergo important functional changes during epileptogenesis (neurogenesis, mossy fiber sprouting, increased excitation; Pitkänen and Lukasiuk, 2011).

To date, three studies have investigated differential expression of miRNAs in the DG during the epileptogenesis and the chronic phase of epilepsy in rats (Bot et al., 2013; Gorter et al., 2014; Roncon et al., 2015). Each of these studies used a different method to trigger epileptogenesis: focal electrical stimulation of the lateral nucleus of the amygdala (Bot et al., 2013); focal electrical stimulation of the angular bundle, a major afferent pathway from the entorhinal cortex to the hippocampus (Gorter et al., 2014); and the systemically administered chemoconvulsant pilocarpine (Roncon et al., 2015). Interestingly, all these models imply a key involvement of the DG in the development of epilepsy but via a different epileptogenic insult: direct activation in the case of angular bundle stimulation, indirect in amygdala stimulation, and a widespread brain activation in the case of pilocarpine (Peng and Houser, 2005).

Here, we aimed to overcome some of the limitations related to individual studies by combining these three studies in a meta-analysis, the aim being to increase the

This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license, which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

Table 1. Datasets included in the meta-analysis

		Bot et al.	Gorter et al.	Dingledine	Dingledine	Dingledine
	Roncon et al. (2015)	(2013)	(2014)	et al. (2017)	et al. (2017)	et al. (2017)
GEO ID	-	GSE49849	-	GSE47752	GSE47752	GSE47752
Rat model	Pilocarpine	Amygdala stimulation	Angular bundle stimulation	Pilocarpine	SSSE	Kainate
Sample count epileptogenesis cases:control	5:5	5:5	8:10	6:6	4:5	6:6
Sample count chronic stage cases:control	5:4	5:5	6:10	-	-	-
Platform	Rat miRNA MicroArray kit, Agilent Technologies	miRCURY LNA microRNA Array7th, Exiqon services	miRCURY LNA microRNA Array 6th, Exiqon services	GeneChip Rat Genome 230 2.0 Array, Affymetrix	GeneChip Rat Genome 230 2.0 Array, Affymetrix	GeneChip Rat Genome 230 2.0 Array, Affymetrix
miRNA/gene expression data	miRNA	miRNA and gene expression	miRNA	Gene expression	Gene expression	Gene expression
Tissue collection	Laser- microdissected DG	Handily dissected DG	Handily dissected DG	Laser- microdissected DG	Laser- microdissected DG	Laser- microdissected DG

statistical power for detecting differentially expressed miRNAs while accounting for study heterogeneity, ultimately leading to more robust and accurate predictions of dysregulated downstream pathways (Ramasamy et al., 2008; Yang et al., 2014). Moreover, this approach offers the opportunity to identify miRNA changes that are independent of the model of epilepsy, i.e., more likely to be disease rather than model related. Our meta-analysis was performed at two time points in the "natural history" of the experimental disease: epileptogenesis and the chronic phase of epilepsy.

Materials and Methods

Inclusion criteria and study design

We collected datasets for meta-analysis based on available genome-wide expression profiles of miRNAs from the DG from epileptic and control hippocampi during epileptogenesis and chronic epilepsy. To assist the functional inference of differentially expressed miRNAs we analyzed publicly available gene expression data obtained from the DG of epileptic rodents.

To identify relevant studies, we first undertook a systematic search to identify all published studies of miRNA expression levels and/or gene expression between cases (epileptic) and controls, in the DG of animal models of epilepsy. We conducted a PubMed search based on the string: "(microRNA OR miRNA) AND (dentate gyrus OR dentate cells OR granule cells) AND epilepsy." miRNAs or genes expression profiles data obtained from the whole hippocampi or different brain regions to the DG were not included in the meta-analysis. This search and inclusion criteria identified only three relevant articles (Bot et al., 2013; Gorter et al., 2014; Roncon et al., 2015).

Time points for each stage of epileptogenesis and for the chronic phase of epilepsy have not been standardized. For the purposes of this study, we considered 7-8 d after SE as the "epileptogenesis phase" and more than two months after SE as the "chronic phase."

The following models were used in the three relevant papers and considered for the meta-analysis. (1) Pilocarpine model: a microarray study based on the investigation of miRNAs differentially expressed in the lasermicrodissected GC layer of the DG of the hippocampi of pilocarpine-induced epileptic rats and matched controls (n = 4), killed during the late phase of latency, 8 d after SE (n = 5) and in the chronic stage, 50 d after the first spontaneous seizure (n = 5; Roncon et al., 2015). (2) Amygdala stimulation: a microarray study focused on the differential expression of miRNAs and genes in handdissected DG in the amygdala stimulation rat model during the phase of epileptogenesis, 7 d after the stimulation (n = 5), in the chronic stage, 60 d after the stimulation (n = 5)= 5) and controls (n = 5; Bot et al., 2013). (3) Perforant path stimulation: a microarray miRNA study based on the perforant path stimulation rat model of epilepsy. We analyzed DG samples obtained from stimulated and control rat hippocampi, during latency (8 d after SE; n = 8) and three to four months after the stimulation for the chronic stage (n = 6) and controls (n = 10; Gorter et al., 2014). The datasets considered for the meta-analysis are summarized in Table 1.

Power calculation

The statistical power of cases and controls for each individual model was calculated using pooled SD of each expressed miRNA. Power to detect miRNAs differentially expressed at multiple fold changes (1, 1.5, 2, 2.5, 3, 3.5, and 4) were calculated, considering miRNA expression variability ranging from 20th, 40th, 60th, and 80th to 100th percentile of the respective SD profiles per model (Fig. 1). Power calculations were performed using R bioconductor package 'pwr' version 1.2.

Data processing

From each identified study the following information was extracted: platform, number of cases and controls, and miRNA expression data at different time points of the disease. When available, GEO accession number and gene expression data were extracted (Table 1).

Data transformation

Since different platforms had been used to generate miRNA expression values, a linear transformation approach was applied to each miRNA using a Z-score transformation according to the formula:

Z-score =
$$\frac{Xi - X}{\delta}$$

Where Xi is the normalized intensity data for each miRNA, X is the average normalized miRNA intensity within a single study, and δ is the SD of cases and controls within respective studies.

Effect size estimation

A meta-analysis is "a technique for quantitatively combining and integrating the results of multiple studies on a given topic" (Polit and Beck, 2004). Thus, a key aspect of meta-analysis is to measure differences and direction of change from quantitative research studies (Polit and Beck, 2004; Berben et al., 2012). A common metric used to provide this important information is the effect size calculation. Accordingly, to give a statistical expression of the magnitude of the difference between groups (i.e., epileptogenesis vs controls and chronic stage vs controls) in regard of miRNAs expression, we estimated the effect size of each individual study defined as the standardized mean difference (SMD) between cases and controls. The SMD has been calculated using the Hedge's method with the following formula:

$$g = \frac{X_1 - X_2}{\delta}$$

Where X_1 is the mean of cases, X_2 the mean of the control group, and δ is the SD.

Statistical heterogeneity

Different animal models, tissue collection methods, and platforms were used to generate the datasets. This makes it difficult to directly compare the data, the risk being of skewing comparison results, reducing the reliability of measurements of individual miRNA expression changes (Yang et al., 2014). Statistical heterogeneity was assessed using Cochrane meta-regression approach calculated by Q test, I² statistics, and Tau² statistics. These measures were applied at each dataset to assess the overall heterogeneity (Higgins et al., 2003; Ioannidis et al., 2007). To test the total variance of each miRNA within the studies, the Cochran Q test have been run, according to the formula:

$$Q = k(k - 1) \frac{\sum_{T=1}^{k} \left(xT - \frac{N}{k} \right)^2}{\sum_{i=1}^{b} xT(k - xT)}$$

Where k is the number of the studies included in the meta-analysis, T is the number of variables observed, b is the number of miRNAs included in the test, and N is the total number. A Benjamini-Hochberg (BH) adjusted p < 0.05 was considered statistically significant for the Co-chrane Q test.

Furthermore, I² statistic has been employed to describe the percentage of the variability in the effect size estimates, following the formula:

$$I^2 = \left(\frac{Q - df}{Q}\right) * 100\%$$

Where Q is the value derived from the Cochran Q test and df are the degrees of freedom. Tentatively, I^2 statistic can be considered as an indicator of heterogeneity, where low, moderate, and high heterogeneity corresponds to I^2 values of 0–0.3, 0.3–0.7, and 0.7–1, respectively (Higgins et al., 2003).

Finally, to estimate the variance across studies the Tau² has been applied:

$$Tau^2 = \frac{Q - df}{C}$$

Where Q is the value derived from the Cochran Q test, df are the degrees of freedom and C is a scaling factor which takes into consideration that the Q value is the weighted sum of squares.

Meta-analysis

The presence of statistical heterogeneity among the studies led us to use a random-effect model for the meta-analysis rather than a fixed-effect model. The pooled effect size (PES) for each miRNA was obtained applying the random effect size model based on the DerSimonian and Laird method (DerSimonian and Laird, 1986; DerSimonian and Kacker, 2007). We generated one forest plot for each miRNA in both the epileptogenesis and the chronic phase to depict the SMD along with its 95% confidence interval (95% CI) for individual studies as well as the pooled MD from the meta-analysis.

miRNAs:mRNAs inverse-fold change

miRNAs predominantly act by repressing their target genes by decreasing target mRNA levels (Bartel, 2004; Guo et al., 2010). Therefore, we investigated the correlation of mRNAs predicted targets expression in the available transcriptomic datasets.

To predict miRNA target genes, the miRNA-target interactions were analyzed with the web-based tool miRwalk (Dweep et al., 2011; Dweep et al., 2014). The rule that the 5' region of miRNA from nucleotides 2-8 ("seed region") has importance in targeting, is commonly accepted as the canonical mechanism by which miRNAs complementary convey functional binding to mRNA targets (Jinek and Doudna, 2009). However, despite the importance of the seed region, the 3' end of a mRNA also contributes to the binding in $\sim 2\%$ of all preferentially conserved sites (Grimson et al., 2007; Shkumatava et al., 2009). In addition, some validated miRNAs have a binding site that exhibits few continuous base pairs in the control region (Shin et al., 2010). Thus, to figure out the complete mechanism of miRNA regulation, we expanded the miRNAs-binding site prediction within the 3', 5' untranslated regions (UTRs), and the seed sequence, with a minimum seed length of seven nucleotides. Furthermore, to exclude overprediction we applied a comparative analysis by six prediction programs: miRMap, RNA22, mi-Randa, RNAhybrid, PICTAR2, and Targetscan. Following this approach, a candidate mRNA target has to be iden-

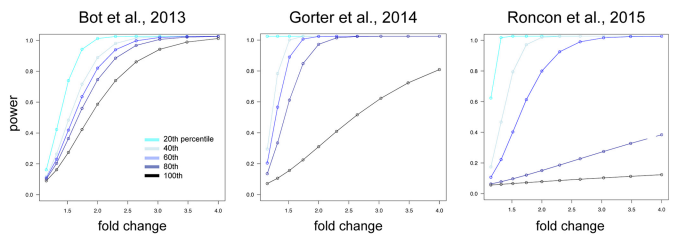


Figure 1. Power calculation. Power calculation is plotted as the power (*y*-axis) to detect a miRNA with fold change (*x*-axis) according to the percentile of the ranked SDs for miRNAs for each study. Across all three models, the power to detect miRNA with fold change 2 or less falls below 80% for at least 40% of the miRNAs.

tified by all these programs. As we did for the miRNAs, we conducted PubMed search based on the keywords: "gene expression, epilepsy, dentate gyrus." Two studies were identified that were then included in our analysis (Table 1): Dingledine et al. (2017; GEO repository accession number: GSE47752) and Bot et al. (2013; GEO repository accession number: GSE49850). The first includes gene expression data obtained from laser-microdissected DG of rats that received different SE models of epilepsy: pilocarpine, self-sustained SE (SSSE), and kainite, it also included kindling, not considered in this study not being a post-SE model. We considered in our analysis only those rats killed 10 d after SE that did not develop spontaneous seizures, as the best time point matching the epileptogenesis phase used for the miRNAs meta-analysis (7-8 d after SE). In Dingledine et al. (2017), the kainate and pilocarpine models were performed in two independent labs, while the SSSE model that was performed in only one lab. For studies undertaken in multiple labs, we combined the p values (obtained from differential expression analysis) from the independent labs using Fisher's method. The second dataset (GSE49850) was obtained from the same amygdala-stimulated rats used for the miRNA analysis.

To investigate miRNA-mRNA interactions, we included only those mRNAs that had inverse relationship to miRNA changes in at least three datasets in the epileptogenesis phase; while for the chronic phase, we considered all predicted mRNAs that presented an inverse relationship to miRNA changes in the amygdala stimulation dataset only.

miRNA functional enrichment analysis

Functional enrichment analysis using gene ontology (GO) and pathways enrichment analysis based on the Kyoto encyclopedia of genes and genomes (KEGG) database were performed using Webgestalt webserver (Zhang et al., 2005; Wang et al., 2013). The enrichment was performed with a hypergeometric test separately for the list of predicted targets based on those miRNAs dysregulated in epileptogenesis and in the chronic stage. Significant canonical pathways maps were selected according to a false discovery rate (FDR) < 5%.

To infer functional relationships between miRNAs identified using meta-analysis, a network of miRNAs based on their ability to target common pathways has been generated. A connection was made between a pair of miRNA, if respective mRNA targets belonged to the same pathways or GO terms that were significantly (FDR < 0.05) enriched for combined miRNA targets. The network was visualizes using Cytoscape version 2.8.2.

Results

Study design

We included in the meta-analysis all published miRNA expression datasets from dissected DG of the hippocampus that compared control (baseline) tissue with tissue from epileptogenesis and chronically epileptic rats (Fig. 2A). Based on these inclusion criteria, we identified three datasets that used different epilepsy models (Table 1): (1) pilocarpine (Roncon et al., 2015), (2) amygdala stimulation (Bot et al., 2013), (3) angular bundle stimulation (Gorter et al., 2014). We first calculated the power of each individual study to detect significant changes in miRNA expression and found that all three individual studies were substantially underpowered to detect modest fold changes (<2.0) in miRNA expression (Fig. 1). Of the total number of miRNAs expressed at any time point in any of the three models, the expression levels of 176 miRNAs were detectable across all three studies (Fig. 2B). The expression values of these 176 miRNAs were then Z-score transformed and metaanalyzed across the three studies.

Estimation of statistical heterogeneity

Meta-regression analyses were performed separately for epileptogenesis and chronic stages of epilepsy for the 176 miRNAs that were expressed in all three datasets (Fig. 2B). There are two models that are commonly used to perform meta-analysis, the fixed effect and the random effects models. The fixed effect model assumes that the

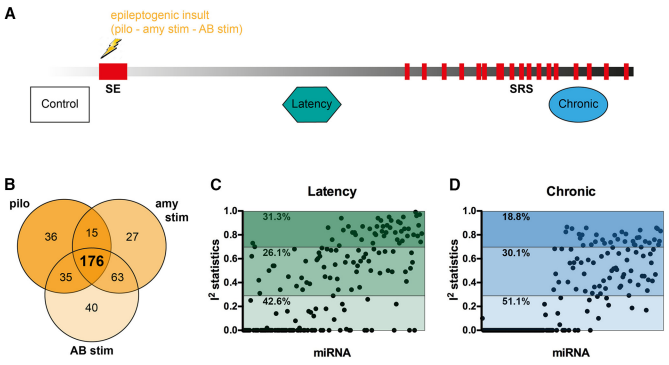


Figure 2. Study design and data preprocessing. *A*, Study design. *B*, Venn diagram showing miRNAs commonly expressed between the three studies included in the meta-analysis. *C*, Statistical heterogeneity estimation. I^2 scores of commonly expressed miRNAs (n = 176) in epileptogenesis and in chronic stage. miRNAs are ordered based on the adjusted *p* value after meta-analysis. $I^2 < 0.3$, low heterogeneity; $0.3 < I^2 > 0.7$, moderate heterogeneity; $I^2 > 0.7$, high heterogeneity. SRS, spontaneous recurrent seizures; pilo, pilocarpine model; amy stim, amygdala stimulation model; AB stim, angular bundle stimulation model.

effect size is the same in all studies, while the random effects analysis assumes that the effect can vary from one study to another. To determine the correct model for this study, we first estimated statistical heterogeneity using the Cochrane's Q test, separately for the epileptogenesis and the chronic phase. This revealed significant heterogeneity between studies at both stages of the disease (BH, adjusted p < 0.05). Next, to assess the proportion of miRNAs that were differentially expressed between studies, during epileptogenesis and the chronic stage separately, we calculated the I² statistics (Higgins et al., 2003). Of the 176 miRNAs measured across the three studies, 26.14% revealed a high level of heterogeneity: 31.25% a moderate level and 42.61% a low rate of heterogeneity during epileptogenesis (Fig. 2C). In the chronic stage, 18.75% displayed high, 30.11% moderate, and 51.14% low level heterogeneity (Fig. 2C). Collectively, these observations favor the use of the random effects model.

Differential expression of miRNAs in the epileptic DG

Using a random effects meta-analysis of miRNA changes in the three models of epileptogenesis and adopting a stringent correction for multiple testing to minimize false positives (Bonferroni adjusted p < 0.05), we identified 26 and 5 differentially expressed miRNAs between control and latency and between control and chronic epilepsy, respectively. Full results of all these miRNAs including PES estimations, I², Tau² and *p* values are shown in Tables 2, 3. Forest plots for selected miRNA are shown in Figure 3. Comparing these results with those

presented in each original studies that have been metaanalyzed here, our meta-analysis identified 11 miRNAs differentially expressed in epileptogenesis compared to control and two miRNAs (i.e., miR-324-3p and miR-130a-3p) in the chronic stage of epilepsy that were not identified as significantly differentially expressed in any of the individual studies (Tables 2, 3, miRNAs highlighted in bold). The datasets employed in this meta-analysis do not allow a precise evaluation of the abundance of expression of these 26 plus five miRNAs under control conditions, but a relative abundance estimate based on the internal standards employed in each study suggests that almost all are expressed at medium to high abundance in the control DG (only miR-212-5p was expressed at relatively low levels but upregulated during epileptogenesis).

Relationship between miRNAs and mRNAs expression changes

Previous transcriptomic studies in epilepsy models have revealed dysregulation of many genes in the different phases of the experimental disease (Lukasiuk and Pitkänen, 2004). It can be hypothesized that differentially expressed miRNAs may contribute to these alterations (upregulated miRNAs may downregulate their mRNA targets whereas downregulation of miRNAs may allow upregulation of their mRNA targets). To assess the role of differentially expressed miRNAs during the course of epilepsy, that is, to infer their mRNA regulatory targets, we explored the potential regulatory relationship between miRNA and mRNA changes.

	ES	Nominal	Bonferroni adjusted			
miRNA	estimation	p value	p value	Q statistics	I ² statistics	Tau ² statistics
miR-212-3p	1.70	6.21 ⁻¹⁶	1.10 ⁻¹³	1.59	0.00	0.00
miR-7a-5p	-1.60	9.26 ⁻¹²	1.64 ⁻⁰⁹	0.82	0.00	0.00
miR-33-5p	-1.57	9.90 ⁻¹²	1.75 ⁻⁰⁹	0.22	0.00	0.00
miR-139-5p	-1.53	8.34 ⁻¹¹	1.48 ⁻⁰⁸	0.29	0.00	0.00
miR-344b-2-3p	1.25	5.78^{-10}	1.02 ⁻⁰⁷	1.57	0.00	0.00
miR-3573-3p	-1.53	1.32^{-09}	2.33 ⁻⁰⁷	1.38	0.00	0.00
miR-551b-3p	-1.51	2.16^{-09}	3.83^{-07}	0.51	0.00	0.00
miR-146a-5p	1.75	3.00^{-09}	5.31 ⁻⁰⁷	2.66	0.25	0.25
miR-132-3p	1.85	1.46 ⁻⁰⁸	2.58^{-06}	5.28	0.62	0.24
let-7b-3p	-1.34	2.46^{-07}	4.36 ⁻⁰⁵	7.40	0.73	0.42
miR-212-5p	1.36	4.58^{-07}	8.11 ⁻⁰⁵	0.36	0.00	0.00
let-7d-3p	-1.42	9.57^{-07}	0.0002	6.75	0.70	0.38
miR-667-3p	-1.37	1.01^{-06}	0.0002	1.87	0.00	0.00
miR-138-5p	-1.40	1.34 ⁻⁰⁶	0.0002	0.59	0.00	0.00
miR-330-3p	-1.37	1.39 ⁻⁰⁶	0.0002	0.17	0.00	0.00
miR-21-5p	1.85	2.65^{-06}	0.0005	3.43	0.42	0.17
miR-29c-5p	-1.65	5.76^{-06}	0.0010	2.91	0.31	0.27
miR-335	-1.38	6.01^{-06}	0.0010	1.41	0.00	0.00
miR-101a-3p	-1.32	9.17^{-06}	0.0016	0.64	0.00	0.00
miR-345-5p	-1.29	2.12 ⁻⁰⁵	0.0037	1.75	0.00	0.00
miR-92b-3p	-1.31	3.12^{-05}	0.0055	1.08	0.00	0.00
miR-150-5p	-1.32	3.46^{-05}	0.0061	1.94	0.00	0.00
miR-136-3p	-1.22	3.56^{-05}	0.0063	0.66	0.00	0.00
miR-324-5p	-1.31	4.63^{-05}	0.0082	2.05	0.02	0.08
miR-153-3p	-1.23	6.92^{-05}	0.0122	0.21	0.00	0.00
miR-383-5p	-1.54	0.0002	0.0375	4.00	0.50	0.34

miRNAs in italics are upregulated and miRNAs highlighted in bold were not differentially expressed in individual studies. ES, effect size.

First, to predict miRNA target transcripts we used the miRWalk database (Dweep et al., 2011; Dweep et al., 2014), which combines information across six miRNA target prediction programs (miRMap, RNA22, miRanda, RNAhybrid, PITCAR2 and Targetscan). As expected, this analysis identified a very large number of predicted targets but, obviously, the large majority of these may not be expressed in the DG or may not be expressed in a negatively correlated fashion relative to the miRNAs. Therefore, we asked whether the miRNAs that were identified by the meta-analysis as dysregulated in epileptogenesis and in the chronic stage were negatively correlated with changes in DG expression in their predicted mRNA targets. To this end, we took advantage of publicly available gene expression data generated in three separate datasets that investigated mRNA expression changes in the DG of rats from the epilepsy models used in our meta-analysis: pilocarpine (GEO accession: GSE47752; Dingledine et al., 2017), angular bundle stimulation (called SSSE in this database; GEO accession: GSE47752), and amygdala stimulation (Bot et al., 2013; PMID: 24146813). Only the amygdala stimulation mRNA dataset (Bot et al., 2013) was obtained from the same animals employed for obtaining the miRNA dataset and included data on the chronic phase. The other datasets were generated by separate research groups on a separate group of animals, the experimental procedures were slightly different from those used in the corresponding miRNAs studies (Gorter et al., 2014; Roncon et al., 2015) and included mRNA data for these models related to the epileptogenesis phase only. In addition, Dingledine et al. (2017) also included datasets for another SE model (kainate) that we also considered in our analysis.

Analysis of the epileptogenesis data identified inverse relationship, based on significant fold changes (gene FDR < 0.1), between 22 (of 26) miRNAs and 122 unique predicted gene targets in at least three of the four epilepsy models of the Dingledine dataset (Dingledine et al., 2017) that we considered in this study (Table 4). The mRNAs encoding the mitogen-activated protein kinase kinase ki-

miRNA miR-652-3p miR-551b-3p miR-324-3p miR-130a-3p	ES estimation -1.56 -1.52 -1.40 -1.35	Nominal p value 5.48 ⁻⁰⁹ 3.20 ⁻⁰⁶ 3.13 ⁻⁰⁵ 6.10 ⁻⁰⁵	Bonferroni adjusted <i>p</i> value 9.65 ⁻⁰⁷ 0.0006 0.0055 0.0107	Q statistics 2.1 1.4 0.9 0.8	l ² statistics 0.1 0.0 0.0 0.0	Tau ² statistics 0.1 0.0 0.0 0.0
miR-130a-3p miR-148b-3p	-1.35 -1.56	0.0002	0.0317	0.8	0.0	0.0

All miRNAs are downregulated, miRNAs highlighted in bold were not differentially expressed in individual studies. ES, effect size.

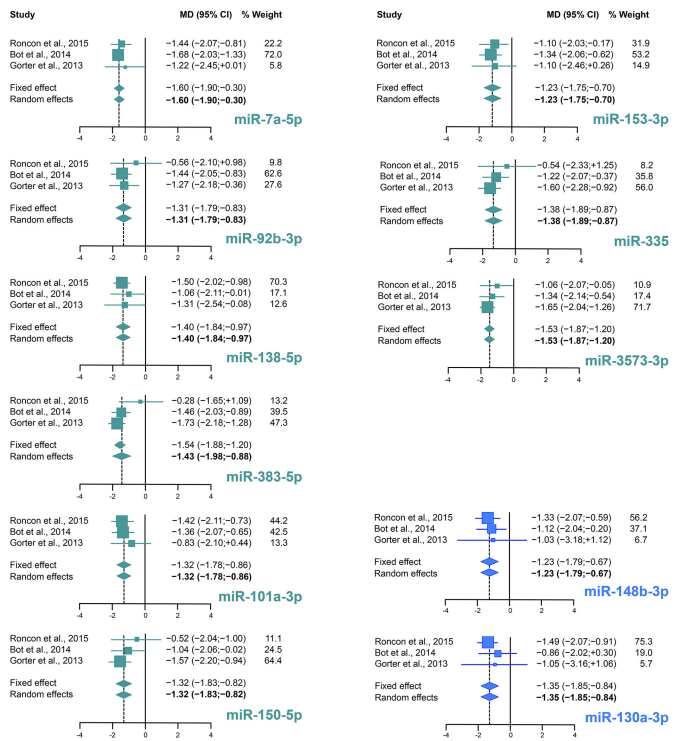


Figure 3. Forest plots of selected miRNA. Forest plots for miR-7a-5p, miR-92b-3p, miR-101a-3p, miR-138-5p, miR-150-3p, miR-153-3p, miR-335, miR-383-3p, and miR-3573-3p are shown for the phase of epileptogenesis, and miR-130a-3p and miR-148b-3p for the chronic stage. For each miRNA, the effect size of the individual studies is reported as MD and 95% CI. The % weight refers to random effects analysis. Individual effect sizes are represented by colored boxes (green for epileptogenesis and blue for the chronic period) and 95% CI are denoted by black lines. The combined effect sizes are represented by diamonds, where diamond width correspond to the 95% CI bounds; boxes and diamonds size is proportional to effect size estimation precision. For each miRNA, the weight of the dataset in the combined analysis has been reported in percentage.

nase 4 (*Map3k4*) and the enhancer of zeste 2 polycomb repressive complex 2 subunit (*Ezh2*) were predicted targets and had inverse expression relative to four miRNAs,

i.e., miR-92b-3p, miR-101a-3p, miR-153-3p, and miR-3575-3p for *Map3k4* and miR-92b-3p, miR-101a-3p, miR-138-5p, and miR-153-3p for *Ezh2*; synapsin type 2 was

Table 4. miRNA-mRNA fold changes inverse correlation in epileptogenesis

	microRNA da	ata			mRNA targets of respective microRNA (FDR $<$ 0.1)								
		Perforant path	Amygdala		mRNA gene	,		Perfor	ant path	Amygo	lala	Kainic	
miRNA name	Pilocarpine	stimulation	stimulation		name	Pilocar	pine	stimula	ation	stimula	ation	acid	
				Meta-analysis		50	Adjusted	50	Adjusted	50	Adjusted	50	Adjuste
miR-383-5p	miRNA FC -0.50	-0.35	-0.75	<i>p</i> value 6.11 ⁻⁰⁵	Rab32	FC 0.74	<i>p</i> value 0.01	FC 0.66	<i>p</i> value 0.01	FC 0.28	<i>p</i> value 0.03	FC 0.48	<i>p</i> value 0.06
min-303-3p	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Cyb561	1.07	4.62^{-05}	0.53	0.01	0.28	1.41 ⁻⁰³	0.48	0.08
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Stk40	0.61	0.05	0.58	0.02	0.35	0.04	0.69	0.04
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Ptpn5	0.59	1.68 ⁻⁰³	0.68	0.02	0.42	0.03	0.72	4.39 ⁻⁰⁴
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Tyms	0.41	0.01	0.81	0.03	0.27	0.06	1.10	3.74^{-04}
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Ugt1a5	0.18	0.06	0.32	0.08	0.69	7.29 ⁻⁰⁴	0.01	0.97
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Aif1	0.91	0.04	0.54	0.25	0.70	0.03	0.97	0.02
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Smad3	0.51	0.06	0.19	0.46	0.45	0.03	0.43	0.08
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Rac2	0.44	0.03	0.39	0.31	0.98	0.01	0.61	0.11
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Lcmt1	0.29	0.04	0.25	0.25	0.32	0.02	0.14	0.48
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Mtmr11	0.95	3.62^{-04}	1.86	0.01	0.05	0.87	1.13	4.82^{-05}
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵	Nnat	1.09	0.01	0.86	0.07	0.15	0.59	0.63	0.30
	-0.50	-0.35	-0.75	6.11 ⁻⁰⁵ 7.95 ⁻⁰⁴	Casp3	0.24	0.07	0.40	0.05	0.27	0.37	0.25	0.40
miR-153-3p	-0.32 -0.32	-0.42 -0.42	-0.22 -0.22	7.95 ⁻⁰⁴	Arhgap17	0.30 0.67	0.02 0.07	0.88 0.79	0.01 0.01	0.43 0.69	0.03 0.01	0.33 0.29	0.08 0.67
	-0.32	-0.42	-0.22	7.95 7.95 ⁻⁰⁴	Mgst1 Wls	1.45	0.07 7.49 ⁻⁰⁵	0.79	0.01	0.69	0.01	0.29 1.75	0.67 1.52 ⁻⁰⁵
	-0.32	-0.42	-0.22	7.95 ⁻⁰⁴	Map3k4	0.09	0.59	0.53	0.03	0.29	0.07	0.02	1.00
	-0.32	-0.42	-0.22	7.95 ⁻⁰⁴	Ezh2	0.05	0.33	0.30	0.07	0.23	0.05	0.02	0.78
	-0.32	-0.42	-0.22	7.95 ⁻⁰⁴	Man2b1	0.10	0.05	0.19	0.63	0.34	0.03	-0.16	0.87
	-0.32	-0.42	-0.22	7.95 ⁻⁰⁴	Zfp521	1.23	1.92^{-04}	0.74	0.00	0.23	0.22	0.59	3.07 ⁻⁰³
miR-324-5p	-0.09	-0.60	-0.29	8.55 ⁻⁰⁶	Tyrobp	1.26	1.01^{-04}	0.70	0.26	1.40	4.93^{-03}	1.31	0.03
-	-0.09	-0.60	-0.29	8.55 ⁻⁰⁶	Asph	0.45	0.08	0.45	0.03	0.17	0.13	0.05	0.90
	-0.09	-0.60	-0.29	8.55 ⁻⁰⁶	Cyb5r4	0.36	0.02	0.43	0.07	0.14	0.66	0.50	7.35^{-04}
miR-150-5p	-0.45	-0.45	-0.56	6.17^{-05}	E2f1	0.89	1.05^{-03}	0.76	0.01	0.45	0.02	1.11	3.99^{-05}
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Cyb561	1.07	4.62^{-05}	0.53	0.02	0.66	1.41^{-03}	0.45	0.09
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Ppp1r1a	0.35	0.03	0.58	0.02	0.44	0.05	0.73	0.01
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Ptpn5	0.59	1.68^{-03}	0.68	0.02	0.42	0.03	0.72	4.39 ⁻⁰⁴
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Tyms	0.41	0.01	0.81	0.03	0.27	0.06	1.10	3.74^{-04}
	-0.45	-0.45	-0.56	6.17^{-05}	lgsf1	1.03	0.37	0.89	0.04	0.31	0.05	0.50	0.49
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵ 6.17 ⁻⁰⁵	Me3	0.73	0.03	0.61	0.07	0.46	1.41 ⁻⁰³ 7.29 ⁻⁰⁴	1.15	4.17 ⁻⁰⁷
	-0.45 -0.45	-0.45 -0.45	-0.56 -0.56	6.17 ⁻⁰⁵	Ugt1a5 Slc7a14	0.18 0.91	0.06 3.46 ⁻⁰⁴	0.32 0.42	0.08 0.09	0.69 0.23	0.06	0.01 0.35	0.97 0.17
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Tmod3	0.72	0.01	0.42	0.09	0.20	0.00	0.33	0.07
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Tyrobp	1.26	1.01 ⁻⁰⁴	0.70	0.26	1.40	4.93 ⁻⁰³	1.31	0.03
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Gpnmb	1.54	3.04^{-03}	0.30	0.68	1.45	0.03	0.81	0.22
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Zmiz1	0.48	2.39 ⁻⁰³	0.33	0.14	0.41	0.03	0.49	0.03
	-0.45	-0.45	-0.56	6.17^{-05}	Skap2	0.73	0.09	0.22	0.37	0.59	0.06	0.57	0.06
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Arhgdib	0.27	0.01	0.48	0.34	0.85	0.03	0.49	0.19
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	lck	0.17	0.04	0.28	0.12	0.27	0.10	-0.01	0.88
	-0.45	-0.45	-0.56	6.17^{-05}	Tmem140	0.80	0.06	0.61	0.05	0.07	0.64	-0.25	0.92
	-0.45	-0.45	-0.56	6.17^{-05}	Trh	1.20	0.01	1.31	0.02	0.99	0.34	2.87	0.01
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Col9a1	0.41	0.02	0.91	0.04	0.03	0.92	1.28	2.67^{-04}
	-0.45	-0.45	-0.56	6.17 ⁻⁰⁵	Arpp21	0.11	0.07	0.45	0.09	-0.02	0.91	0.63	0.02
miR-92b-3p	-1.05	-0.72	-0.37	1.70 ⁻⁰⁵	Gadd45a	-0.20	0.16	1.51	0.01	0.79	0.01	1.33	3.91 ⁻⁰⁴
	-1.05	-0.72	-0.37	1.70^{-05}	Map3k4	0.09	0.59	0.53	0.03	0.29	0.07	0.02	1.00
	-1.05	-0.72	-0.37	1.70 ⁻⁰⁵ 1.70 ⁻⁰⁵	Kcnh2	0.60	0.07	0.63	0.04	0.39	0.09	0.26	0.38
	-1.05 -1.05	-0.72 -0.72	-0.37 -0.37	1.70 ⁻⁰⁵	Ezh2 Wnt10a	0.16 0.58	0.27 1.89 ⁻⁰³	0.40 0.81	0.07 0.07	0.31 0.46	0.05 0.01	0.04 1.69	0.78 0.01
	-1.05	-0.72	-0.37 -0.37	1.70 ⁻⁰⁵	Zmiz1	0.58	2.39 ⁻⁰³	0.81	0.07	0.46	0.01	0.49	0.01
	-1.05	-0.72	-0.37	1.70 ⁻⁰⁵	lck	0.40	0.04	0.33	0.14	0.41	0.03	-0.01	0.88
	-1.05	-0.72	-0.37	1.70 ⁻⁰⁵	Zfp521	1.23	1.92 ⁻⁰⁴	0.20	0.12	0.27	0.10	0.59	3.07 ⁻⁰³
miR-345-5p	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Inpp4b	0.80	0.10	1.15	2.69^{-03}	0.20	0.22	0.33	0.13
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Arhgap17	0.30	0.02	0.88	0.01	0.43	0.03	0.33	0.08
	-0.23	-0.25	-0.12	1.46^{-06}	Gadd45a	-0.20	0.16	1.51	0.01	0.79	0.01	1.33	3.91 ⁻⁰⁴
	-0.23	-0.25	-0.12	1.46^{-06}	Gcnt1	1.45	3.63^{-05}	1.61	0.06	0.48	0.01	1.87	4.39^{-04}
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	LOC500956	0.34	0.20	0.50	0.07	0.27	0.02	0.12	0.91
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Cd74	1.10	0.04	1.27	0.07	2.81	0.01	1.24	0.11
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Wnt10a	0.58	1.89^{-03}	0.81	0.07	0.46	0.01	1.69	0.01
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Skap2	0.73	0.09	0.22	0.37	0.59	0.06	0.57	0.06
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Ss18	0.41	0.02	0.51	0.12	0.58	0.02	0.76	3.58^{-04}
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Slfn13	0.46	0.04	0.41	0.18	1.53	2.69 ⁻⁰³	0.40	0.12
	-0.23	-0.25	-0.12	1.46 ⁻⁰⁶	Rnd3	0.87	0.01	0.60	0.07	0.13	0.56	0.36	0.13
10 404 F	-0.23	-0.25	-0.12	1.46^{-06}	Tmem140	0.80	0.06	0.61	0.05	0.07	0.64	-0.25	0.92
	-0.68	-0.71	-0.19	5.15^{-06}	Map3k4	0.09	0.59	0.53	0.03	0.29	0.07	0.02	1.00
miR-101a-3p	-0.68	-0.71	-0.19	5.15 ⁻⁰⁶	Rin2	0.57	0.06	0.43	0.05	0.44	0.03	-0.06	0.95

Table 4. Continued

	microRNA da	ata	mRNA targets of respective microRNA (FDR $<$ 0.1)										
		Perforant path	Amygdala		mRNA gene	,		Perfora	int path	Amygd	lala	Kainic	
miRNA name	Pilocarpine	stimulation	stimulation		name	Pilocar		stimula		stimula		acid	
	miRNA FC			Meta-analysis		50	Adjusted	50	Adjusted	50	Adjusted	50	Adjuste
	-0.68	-0.71	-0.19	p value 5.15 ⁻⁰⁶	Ezh2	FC 0.16	p value 0.27	FC 0.40	<i>p</i> value 0.07	FC 0.31	<i>p</i> value 0.05	FC 0.04	p value 0.78
	-0.68	-0.71	-0.19	5.15 ⁻⁰⁶	Arl4a	0.65	0.02	0.36	0.41	0.40	0.03	0.38	0.09
	-0.68	-0.71	-0.19	5.15 ⁻⁰⁶	Syn2	0.65	0.04	0.55	0.03	0.11	0.21	0.39	0.03
	-0.68	-0.71	-0.19	5.15 ⁻⁰⁶	Nabp1	0.56	0.01	0.81	0.01	0.45	0.11	0.96	0.10
	-0.68	-0.71	-0.19	5.15 ⁻⁰⁶	Arpp21	0.11	0.07	0.45	0.09	-0.02	0.91	0.63	0.02
niR-335	-1.98	-0.39	-0.71	2.47^{-05}	Efr3a	1.28	0.01	1.40	5.22^{-04}	0.49	0.01	1.11	4.17^{-07}
	-1.98	-0.39	-0.71	2.47 ⁻⁰⁵	Rprm	2.68	3.86^{-05}	2.54	8.57^{-04}	1.00	0.01	1.53	5.05^{-04}
	-1.98	-0.39	-0.71	2.47 ⁻⁰⁵	Ackr3	0.90	0.39	2.38	2.69^{-03}	0.55	3.05^{-03}	0.90	0.11
	-1.98	-0.39	-0.71	2.47 ⁻⁰⁵	Gcnt1	1.45	3.63 ⁻⁰⁵	1.61	0.06	0.48	0.01	1.87	4.39 ⁻⁰⁴
	-1.98	-0.39	-0.71	2.47 ⁻⁰⁵	Fcgr2b	1.87	3.62^{-04}	0.99	0.20	1.52	0.01	1.39	0.01
	-1.98	-0.39	-0.71	2.47 ⁻⁰⁵	Vim	1.74	0.01	1.01	0.28	0.72	0.05	0.59	0.88
	-1.98	-0.39	-0.71	2.47^{-05}	Arl11	0.86	0.01	0.08	0.75	0.72	0.01	0.71	0.07
	-1.98	-0.39	-0.71	2.47^{-05}	Epsti1	0.92	0.03	0.47	0.35	0.71	0.02	0.61	0.02
·D 00 5	-1.98	-0.39	-0.71	2.47 ⁻⁰⁵ 1.48 ⁻⁰⁷	Pycard	0.87	0.01	0.47	0.13	0.47	0.09	0.84	0.01 3.99 ⁻⁰⁶
niR-29c-5p	-0.59	-0.41	-0.14	1.48 ⁻⁰⁷	E2f1	0.89	1.05 ⁻⁰³	0.76	0.01	0.45	0.02	1.11	
	-0.59	-0.41	-0.14	1.48 ⁻⁰⁷	Tmem176b	0.75	0.48	0.67	0.05	0.97	0.01 4.00 ⁻⁰³	1.12	0.02
D 220 2-	-0.59	-0.41	-0.14	1.48 ⁻⁰⁶	Sh3bgrl3	0.77	0.01	0.49	0.13	0.48		0.70	0.01 4.17 ⁻⁰⁷
niR-330-3p	-0.99 -0.99	-0.40	-0.32 -0.32	1.94 ⁰⁰ 1.94 ⁻⁰⁶	Efr3a Sorino2	1.28	0.01 1.52 ⁻⁰⁷	1.40	0.00	0.49	0.01 7.30 ⁻⁰⁴	1.11	4.17 ⁻⁰⁷ 4.02 ⁻⁰⁶
	-0.99 -0.99	-0.40 -0.40	-0.32 -0.32	1.94 ⁰⁰ 1.94 ⁻⁰⁶	Serinc2 Tmem176b	1.16 0.75	0.48	1.47 0.67	0.01 0.05	1.03 0.97	0.01	1.43 1.12	4.02 ⁰⁰
	-0.99 -0.99	-0.40	-0.32 -0.32	1.94 ⁻⁰⁶	Arl11	0.75	0.48	0.67	0.05	0.97	0.01	0.71	0.02
	-0.99 -0.99	-0.40	-0.32 -0.32	1.94 1.94 ⁻⁰⁶	Pycard	0.86	0.01	0.08	0.75	0.72	0.01	0.71	0.07
	-0.99	-0.40	-0.32	1.94 ⁻⁰⁶	Dhrs4	0.20	0.10	0.47	0.13	0.47	0.03	-0.03	0.87
	-0.99	-0.40	-0.32	1.94 ⁻⁰⁶	Cd44	0.20	0.09	0.01	0.99	0.68	0.02	0.18	0.67
	-0.99	-0.40	-0.32	1.94 ⁻⁰⁶	Anks1a	0.72	0.03	0.53	0.05	0.28	0.19	0.17	0.53
	-0.99	-0.40	-0.32	1.94 ⁻⁰⁶	Asph	0.45	0.08	0.45	0.03	0.17	0.13	0.05	0.90
	-0.99	-0.40	-0.32	1.94 ⁻⁰⁶	Cald1	0.30	0.08	0.38	0.09	0.15	0.33	0.07	0.68
niR-138-5p	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Rab32	0.74	0.01	0.66	0.01	0.28	0.03	0.48	0.06
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Tpbg	0.83	0.11	1.71	0.01	0.39	0.01	1.70	0.00
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Map3k14	0.11	0.08	0.77	0.03	0.26	0.07	0.20	0.87
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Tcirg1	0.05	0.82	0.36	0.07	0.46	0.06	-0.10	0.89
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Kank2	0.62	0.02	0.71	0.07	0.59	0.02	2.91 ⁻⁰³	0.95
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Ezh2	0.16	0.27	0.40	0.07	0.31	0.05	0.04	0.78
	-0.48	-0.29	-0.50	3.98^{-08}	Spsb1	-0.08	0.22	0.41	0.09	0.40	0.03	0.05	0.99
	-0.48	-0.29	-0.50	3.98^{-08}	C1qc	1.11	1.37^{-03}	0.42	0.34	0.97	0.08	0.92	0.18
	-0.48	-0.29	-0.50	3.98^{-08}	Sh3bgrl3	0.77	0.01	0.49	0.13	0.48	0.00	0.70	0.01
	-0.48	-0.29	-0.50	3.98^{-08}	Zmiz1	0.48	2.39^{-03}	0.33	0.14	0.41	0.03	0.49	0.03
	-0.48	-0.29	-0.50	3.98^{-08}	Ly86	0.87	0.03	0.40	0.41	0.97	0.03	0.58	0.05
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Fancd2os	0.66	0.04	0.36	0.18	0.28	0.04	0.85	4.31 ⁻⁰³
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Slc20a1	0.41	0.02	0.14	0.52	0.45	0.00	0.27	0.09
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Rassf5	0.36	0.08	0.00	1.00	0.32	0.05	0.46	0.43
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Cd44	0.11	0.09	0.01	0.99	0.68	0.02	0.18	0.67
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Zfp521	1.23	1.92^{-04}	0.74	0.01	0.23	0.22	0.59	3.07 ⁻⁰³
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸	Anks1a	0.72	0.03	0.53	0.05	0.28	0.19	0.17	0.53
	-0.48	-0.29	-0.50	3.98^{-08}	Nnat	1.09	0.01	0.86	0.07	0.15	0.59	0.63	0.30
	-0.48	-0.29	-0.50	3.98^{-08}	Htatip2	0.70	0.01	0.37	0.07	-0.02	0.96	0.79	7.35 ⁻⁰⁴
	-0.48	-0.29	-0.50	3.98^{-08}	Tnfsf9	0.58	0.03	0.78	0.03	0.06	0.63	0.97	1.99 ⁻⁰⁴
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸ 3.98 ⁻⁰⁸	Nabp1	0.56	0.01	0.81	0.01	0.45	0.11	0.96	0.10
	-0.48	-0.29	-0.50	3.98 ⁻⁰⁸ 3.98 ⁻⁰⁸	Numbl Cold1	0.37	0.06	0.40	0.03	0.20	0.14	0.43	0.32 0.68
	-0.48 -0.48	-0.29 -0.29	-0.50 -0.50	3.98 ⁻⁰⁸	Cald1 Arop21	0.30 0.11	0.08 0.07	0.38 0.45	0.09 0.09	0.15 -0.02	0.33 0.91	0.07 0.63	0.68
niR-667-3p	-0.48 -0.34	-0.29 -0.20	-0.50 -0.49	3.98 1.62 ⁻⁰⁷	Arpp21 Mdm1	0.11	0.07	0.45	0.09 2.69 ⁻⁰³	-0.02 0.30	0.91	0.63	0.02
-op	-0.34 -0.34	-0.20	-0.49 -0.49	1.62 ⁻⁰⁷	Inpp4b	0.80	0.04	1.15	2.69 ⁻⁰³	0.30	0.03	0.29	0.07
	-0.34 -0.34	-0.20	-0.49 -0.49	1.62^{-07}	трр46 Тјр2	0.80	0.10	0.78	0.02	0.31	0.07	0.24	0.13
	-0.34 -0.34	-0.20	-0.49 -0.49	1.62 ⁻⁰⁷	1jp2 Serping1	1.19	0.15 3.39 ⁻⁰³	1.22	0.02	0.42 2.12	0.00	1.15	0.56
	-0.34 -0.34	-0.20	-0.49 -0.49	1.62^{-07}	Tmem176b	0.75	0.48	0.67	0.02	0.97	0.00	1.15	0.15
	-0.34	-0.20	-0.49	1.62^{-07}	LOC500956	0.34	0.40	0.50	0.03	0.37	0.01	0.12	0.02
	-0.34	-0.20	-0.49	1.62^{-07}	Chi3l1	1.14	0.20	0.02	0.97	0.27	0.02	0.33	0.82
	-0.34	-0.20	-0.49	1.62^{-07}	Laptm5	0.69	0.03	0.31	0.43	1.22	0.00	0.71	0.02
	-0.34	-0.20	-0.49	1.62^{-07}	lfi30	0.62	0.10	0.11	0.73	0.67	0.07	0.26	0.65
	-0.34	-0.20	-0.49	1.62^{-07}	Fcgr1a	0.33	0.04	0.33	0.26	0.61	0.03	0.33	0.57
	-0.34	-0.20	-0.49	1.62^{-07}	Tmem140	0.80	0.04	0.61	0.05	0.07	0.64	-0.25	0.92
niR-212-5p	2.55	0.27	0.28	1.15 ⁻⁰⁵	Rasd2	-0.59	1.03 ⁻⁰³	-1.24	1.19 ⁻⁰³	-0.87	0.00	-0.58	3.80 ⁻⁰³
1-	2.55	0.27	0.28	1.15 ⁻⁰⁵	Gabrd	-0.53	0.01	-0.59	0.01	-0.55	0.01	-0.71	0.03
	2.55	0.27	0.28	1.15 ⁻⁰⁵	Hpca	-0.74	0.02	-0.64	0.01	-0.22	0.02	-0.40	0.09
					(Continued			-				-	

	microRNA da	ata	mRNA targets of respective microRNA (FDR < 0.1)										
		Perforant path	Amygdala	mRNA gene			Perforant path				ala	Kainic	
miRNA name	Pilocarpine	stimulation	stimulation		name	Pilocar		stimula	ation	stimula		acid	
	miRNA FC			Meta-analysis p value		FC	Adjusted p value	FC	Adjusted p value	FC	Adjusted p value	FC	Adjusteo p value
	2.55	0.27	0.28	1.15 ⁻⁰⁵	C1ql3	0.14	0.57	-0.47	0.05	-0.38	0.00	-0.61	0.38
	2.55	0.27	0.28	1.15^{-05}	Fat1	-1.05	1.24^{-03}	-1.01	0.07	-0.19	0.08	-1.55	1.24^{-03}
let-7b-3p	-0.80	-0.41	-5.75	8.36 ⁻⁰⁶	C1r	0.87	8.05^{-04}	0.79	0.02	0.81	0.01	0.54	0.01
	-0.80	-0.41	-5.75	8.36 ⁻⁰⁶	Gcnt1	1.45	3.63^{-05}	1.61	0.06	0.48	0.01	1.87	4.39^{-04}
	-0.80	-0.41	-5.75	8.36 ⁻⁰⁶	Bdnf	0.45	0.04	0.20	0.69	0.32	0.05	1.17	0.03
	-0.80	-0.41	-5.75	8.36 ⁻⁰⁶	Gpr83	0.59	0.01	0.90	0.06	0.23	0.40	0.43	0.51
	-0.80	-0.41	-5.75	8.36 ⁻⁰⁶	Jup	0.38	0.02	0.45	0.09	0.23	0.29	0.74	1.73^{-03}
miR-132-3p	1.91	0.80	0.10	4.19 ⁻¹⁸	Insig2	-0.37	0.52	-0.81	0.01	-0.34	0.10	-0.15	0.80
miR-146a-5p	4.71	0.37	0.33	4.05 ⁻¹⁰	Mthfd11	-0.94	5.30^{-04}	-0.83	0.01	-0.44	0.01	-1.27	0.04
	4.71	0.37	0.33	4.05 ⁻¹⁰	Plxdc1	-0.28	0.06	-0.73	0.01	-0.33	0.08	-0.70	7.35 ⁻⁰⁴
	4.71	0.37	0.33	4.05 ⁻¹⁰	Htr5b	-0.54	0.01	-1.66	0.03	-0.73	0.00	-1.55	3.87^{-05}
	4.71	0.37	0.33	4.05 ⁻¹⁰	B3galt5	-1.39	7.49^{-05}	-0.83	0.04	-0.43	0.05	-1.93	0.01
	4.71	0.37	0.33	4.05 ⁻¹⁰	Pip5k1b	-0.79	0.17	-0.46	0.05	-0.38	0.06	-1.03	0.02
miR-551b-3p	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Efr3a	1.28	0.01	1.40	5.22 ⁻⁰⁴	0.49	0.01	1.11	4.17 ⁻⁰⁷
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Rprm	2.68	3.86 ⁻⁰⁵	2.54	8.57 ⁻⁰⁴	1.00	0.01	1.53	5.05 ⁻⁰⁴
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Lox	3.11	2.13 ⁻⁰⁹	2.56	2.69^{-03}	1.28	0.00	2.66	6.51 ⁻⁰⁶
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Sox11	1.57	0.01	1.36	0.01	1.23	0.00	1.52	2.72 ⁻⁰³
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Tpbg	0.83	0.11	1.71	0.01	0.39	0.01	1.70	3.06^{-03}
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	C1qc	1.11	1.37 ⁻⁰³	0.42	0.34	0.97	0.08	0.92	0.18
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Syn2	0.65	0.04	0.55	0.03	0.11	0.21	0.39	0.03
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Nabp1	0.56	0.01	0.81	0.01	0.45	0.11	0.96	0.10
	-0.78	-0.85	-0.42	9.09 ⁻¹⁴	Ntm	0.71	0.05	0.37	0.09	-0.22	0.37	-0.32	0.94
miR-3573-3p	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Serping1	1.19	3.39^{-03}	1.22	0.02	2.12	0.00	1.15	0.15
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Map3k4	0.09	0.59	0.53	0.03	0.29	0.07	0.02	1.00
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Col6a3	0.31	0.01	0.79	0.08	0.35	0.06	0.63	0.09
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Plcxd3	1.94	3.86 ⁻⁰⁵	1.58	0.17	0.59	0.01	2.02	0.01
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	S100a10	1.36	1.68^{-03}	0.61	0.50	0.80	0.02	1.09	0.43
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Chi3l1	1.14	0.09	0.02	0.97	0.31	0.08	0.33	0.82
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Rbms1	1.24	1.76^{-03}	0.65	0.17	0.50	0.08	0.59	0.34
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Lgmn	0.41	0.03	0.09	0.77	0.61	0.06	0.22	0.62
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	P2ry6	0.17	0.05	0.12	0.72	0.61	0.02	0.50	0.16
	-1.02	-0.26	-0.93	4.68 ⁻¹⁷	Epb41l4b	0.70	8.37 ⁻⁰⁴	0.58	0.02	0.17	0.46	0.57	2.98 ⁻⁰³
miR-139-5p	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	C1s	1.55	1.42^{-06}	1.15	2.00^{-03}	0.55	0.02	1.34	3.94^{-05}
	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	Tmem176b	0.75	0.48	0.67	0.05	0.97	0.01	1.12	0.02
	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	Slc7a14	0.91	3.46 ⁻⁰⁴	0.42	0.09	0.23	0.06	0.35	0.17
	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	Fcgr2b	1.87	3.62^{-04}	0.99	0.20	1.52	0.01	1.39	0.01
	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	C5ar1	0.44	0.02	0.33	0.31	0.24	0.08	0.43	0.28
	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	Syn2	0.65	0.04	0.55	0.03	0.11	0.21	0.39	0.03
	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	Anks1a	0.72	0.03	0.53	0.05	0.28	0.19	0.17	0.53
	-1.40	-0.77	-0.91	4.40 ⁻¹⁷	Mtmr11	0.95	3.62 ⁻⁰⁴	1.86	0.01	0.05	0.87	1.13	4.82 ⁻⁰⁵
miR-33-5p	-2.40	-0.80	-0.51	4.92 ⁻¹⁹	Runx1	1.08	7.92^{-04}	1.48	2.69^{-03}	0.60	2.69^{-03}	1.22	8.95 ⁻⁰⁴
	-2.40	-0.80	-0.51	4.92^{-19}	Wnt10a	0.58	1.89 ⁻⁰³	0.81	0.07	0.46	0.01	1.69	0.01
	-2.40	-0.80	-0.51	4.92^{-19}	Slc7a14	0.91	3.46^{-04}	0.42	0.09	0.23	0.06	0.35	0.17
	-2.40	-0.80	-0.51	4.92^{-19}	Fcgr2b	1.87	3.62 ⁻⁰⁴	0.99	0.20	1.52	0.01	1.39	0.01
	-2.40	-0.80	-0.51	4.92^{-19}	Ly86	0.87	0.03	0.40	0.41	0.97	0.03	0.58	0.05
B Za Fr	-2.40	-0.80	-0.51	4.92^{-19}	Cfh	1.14	0.04	0.91	0.21	0.70	0.03	0.70	0.11
miR-7a-5p	-0.91	-0.93	-0.51	3.44^{-24}	Mdm1	0.69	0.04	0.76	2.69^{-03}	0.30	0.03	0.29	0.07
	-0.91	-0.93	-0.51	3.44^{-24}	Wls	1.45	7.49 ⁻⁰⁵	0.80	0.01	0.57	0.01	1.75	1.52^{-05}
	-0.91	-0.93	-0.51	3.44^{-24}	Tpbg	0.83	0.11	1.71	0.01	0.39	0.01	1.70	3.06 ⁻⁰³
	-0.91	-0.93	-0.51	3.44^{-24}	Serping1	1.19	3.39 ⁻⁰³	1.22	0.02	2.12	0.00	1.15	0.15
	-0.91	-0.93	-0.51	3.44^{-24}	Map3k14	0.11	0.08	0.77	0.03	0.26	0.07	0.20	0.87
	-0.91	-0.93	-0.51	3.44^{-24}	Tmem176b	0.75	0.48	0.67	0.05	0.97	0.01	1.12	0.02
	-0.91	-0.93	-0.51	3.44^{-24}	Dnah12	0.32	0.41	0.65	0.05	0.22	0.03	-0.35	0.91
	-0.91	-0.93	-0.51	3.44^{-24}	Pafah1b3	0.88	0.04	0.84	0.09	0.28	0.01	0.42	0.79
	-0.91	-0.93	-0.51	3.44^{-24}	S100a4	1.44	2.54 ⁻⁰⁷	0.53	0.17	0.67	0.01	1.05	0.09
	-0.91	-0.93	-0.51	3.44^{-24}	C1qa	1.21	0.05	0.34	0.29	1.27	0.04	1.16	3.03 ⁻⁰³
	-0.91	-0.93	-0.51	3.44^{-24}	Gpnmb	1.54	3.04^{-03}	0.30	0.68	1.45	0.03	0.81	0.22
	-0.91	-0.93	-0.51	3.44^{-24}	Resp18	0.65	1.68 ⁻⁰³	0.38	0.23	0.28	0.09	0.82	0.02
	-0.91	-0.93	-0.51	3.44^{-24}	Cfh	1.14	0.04	0.91	0.21	0.70	0.03	0.70	0.11
	-0.91	-0.93	-0.51	3.44^{-24}	Arhgdib	0.27	0.01	0.48	0.34	0.85	0.03	0.49	0.19
	-0.91	-0.93	-0.51	3.44^{-24}	lck	0.17	0.04	0.28	0.12	0.27	0.10	-0.01	0.88
	-0.91	-0.93	-0.51	3.44^{-24}	Nubpl	1.06	1.77 ⁻⁰⁴	0.88	2.44^{-03}	0.06	0.64	0.77	0.02
	-0.91	-0.93	-0.51	3.44^{-24}	Arpp21	0.11	0.07	0.45	0.09	-0.02	0.91	0.63	0.02
miR-212-3p	1.09	0.88	0.22	1.74^{-47}	Insig2	-0.37	0.52	-0.81	0.01	-0.34	0.10	-0.15	0.80

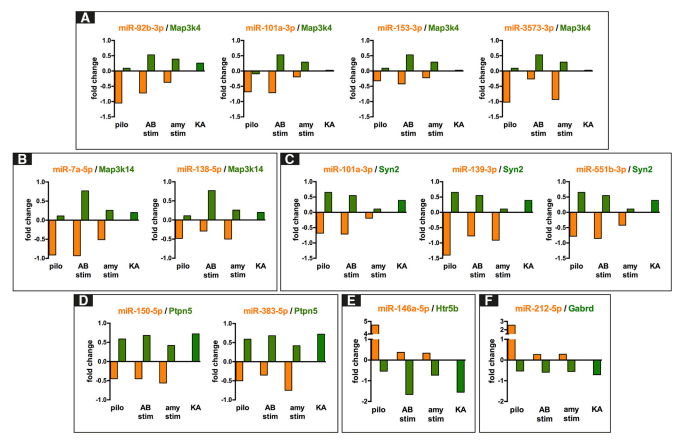


Figure 4. Relationship between selected miRNA and their predicted targets in different model of TLE. All panels show selected miRNAs-mRNA anticorrelation based on miRNAs and mRNAs fold changes in epileptogenesis. *A*, Inverse relationship between four downregulated miRNAs (miR-92b-3p, miR-101a-3p, miR-153-3p, and miR-3573-3p) and the commonly predicted target *Map3k4*. *B*, Inverse relationship between the downregulated miR-138-5p, miR-7a-5p, and the upregulated *Map3k14*. *C*, Inverse relationship between miR-101a-3p, miR-139-3p, miR-551b-3p, and *Syn2*. *D*, Inverse relationship between miR-150-5p, miR-383-5p, and *Ptpn*. *E*, *F*, Examples of the opposite anticorrelation, the upregulated miR-146a-5p with the downregulated *Htr5b* transcript, and the upregulated miR-212-5p and the downregulated *Gabrd* transcript.

inversely correlated to three miRNAs (miR-101a-3p, miR-139-5p, and miR-551b-3p); the mitogen-activated protein kinase kinase kinase 14 (Map3k14) and the protein tyrosine phosphatase, nonreceptor type 5 (Ptpn5) were inversely correlated with two miRNAs, i.e., miR-7a-5p and miR-138-5p for Map3k14; miR-150-5p and miR-383-5p for Ptpn5. All the above transcripts were upregulated during epileptogenesis and predicted targets of miRNAs that were downregulated. In contrast, miR-132-3p, miR-146a-5p, miR-212-3p, and miR-212-5p were upregulated during epileptogenesis. The 5-hydroxytryptamine receptor 5 (Htr5b) and the β -1,3-galactosyltransferase 5 (B3galt5) were predicted targets of miR-146a-5p and were downregulated. The γ -aminobutiric acid receptor subunit δ (Gabrd) was a predicted target of and inversely correlated with miR-212-5p. Finally, we observed that miR-344b-2-3p, let-7d-3p, miR-21-5p, miR-29c-5p, and miR-324-5p were not anticorrelated with any of their predicted mRNA targets. Representative graphs for the anticorrelations between miRNAs and predicted mRNA targets are shown in Figure 4.

The relationship between the changes in expression of miRNAs and their mRNA targets in the chronic stage of epilepsy was analyzed using only the amygdala stimulation

November/December 2017, 4(6) e0152-17.2017

dataset (Bot et al., 2013). We observed negative correlations (based on fold changes) between all five miRNAs that emerged as significantly downregulated from the metaanalysis and 29 unique predicted mRNA targets in the dataset. Five of these 29 anticorrelated mRNAs were predicted targets and had inverse expression relative to two miRNAs and one, the glutamate ionotropic receptor δ -type subunit 2 (*Grid2*), was a predicted target and had inverse expression relative to three miRNAs, namely, miR-130a-3p, miR-148b-3p, and miR-551b-3p (Table 5). Furthermore, interestingly, three mRNA targets, the transmembrane protein 176B (*Tmem176b*), the EFR3 homolog A (*Efr3a*), and the zinc finger, MIZ-type containing 1 (*Zmiz1*) were downregulated in both epileptogenesis and the chronic stage.

Recent evidence supports the notion that miRNAs not only decrease levels of their mRNA targets (Guo et al., 2010), but additionally may have nuclear functions capable of influencing gene expression, and which may be reflected by a correlation between a miRNA and its target gene mRNA levels (Catalanotto et al., 2016). Analysis of the epileptogenesis data revealed significant correlation (gene FDR < 0.1), for 21 (of 26) miRNAs and 77 unique predicted gene targets in at least three of the four epilepsy

Table 5. miRNA-mRNA fold changes inverse correlation at the chronic stage

	Pilocarpine	Angular bundle stimulation	Amygdala stimulation			Amygd Stimula	
				Meta-analysis			
				adjusted	mRNA		adjusted
miRNA name	miRNA FC			p value	gene name	FC	<i>p</i> value
miR-130a-3p	-0.24	-0.32	-0.46	0.0107	Frmd6	0.35	0.0077
	-0.24	-0.32	-0.46	0.0107	Grid2	0.31	0.0702
	-0.24	-0.32	-0.46	0.0107	Necab3	0.43	0.0202
	-0.24	-0.32	-0.46	0.0107	Npepl1	0.22	0.0847
miR-148b-3p	-0.24	-0.29	-0.41	0.0317	C1qa	0.81	0.0559
	-0.24	-0.29	-0.41	0.0317	Ctsz	0.63	0.0773
	-0.24	-0.29	-0.41	0.0317	Flnc	0.53	0.0455
	-0.24	-0.29	-0.41	0.0317	Frmd6	0.35	0.0077
	-0.24	-0.29	-0.41	0.0317	Grid2	0.31	0.0702
	-0.24	-0.29	-0.41	0.0317	Npepl1	0.22	0.0847
	-0.24	-0.29	-0.41	0.0317	Tax1bp3	0.36	0.0371
	-0.24	-0.29	-0.41	0.0317	Tmem176b	0.91	0.0621
miR-324-3p	-0.44	-0.22	-0.27	0.0055	Acss1	0.31	0.0899
	-0.44	-0.22	-0.27	0.0055	Atraid	0.23	0.0936
	-0.44	-0.22	-0.27	0.0055	Cd9	0.46	0.0380
	-0.44	-0.22	-0.27	0.0055	Chi3l1	0.45	0.0455
	-0.44	-0.22	-0.27	0.0055	Csf1r	0.80	0.0773
	-0.44	-0.22	-0.27	0.0055	Ctsb	0.20	0.0918
	-0.44	-0.22	-0.27	0.0055	Gfap	0.93	0.0380
	-0.44	-0.22	-0.27	0.0055	Gsap	0.20	0.0843
	-0.44	-0.22	-0.27	0.0055	Hmox1	0.19	0.0972
	-0.44	-0.22	-0.27	0.0055	Limd2	0.26	0.0817
	-0.44	-0.22	-0.27	0.0055	Mex3b	0.33	0.0217
	-0.44	-0.22	-0.27	0.0055	Osbpl9	0.19	0.0760
	-0.44	-0.22	-0.27	0.0055	Slco2b1	0.46	0.0896
	-0.44	-0.22	-0.27	0.0055	Tmem176b	0.91	0.0621
	-0.44	-0.22	-0.27	0.0055	Zmiz1	0.34	0.0077
miR-551b-3p	-1.30	-0.52	-0.50	0.0006	Csf1r	0.80	0.0773
	-1.30	-0.52	-0.50	0.0006	Efr3a	0.60	0.0027
	-1.30	-0.52	-0.50	0.0006	Entpd2	0.44	0.0518
	-1.30	-0.52	-0.50	0.0006	Grid2	0.31	0.0702
	-1.30	-0.52	-0.50	0.0006	Npc2	0.80	0.0882
	-1.30	-0.52	-0.50	0.0006	Sox11	0.86	0.0455
miR-652-3p	-0.90	-0.34	-0.41	9.65^{-07}	Cd9	0.46	0.0380
	-0.90	-0.34	-0.41	9.65^{-07}	Hsd3b7	0.31	0.0402
	-0.90	-0.34	-0.41	9.65^{-07}	Tmem176a	0.93	0.0559

models of the Dingledine dataset (Dingledine et al., 2017; Table 6). In addition, we found positive correlations between five of the five miRNAs that were downregulated in the chronic period and 39 predicted mRNA targets in the amygdala stimulation dataset (Bot et al., 2013; Table 7). Interestingly, 29 of the mRNAs identified as potential targets in epileptogenesis were inversely correlated to some miRNAs and directly correlated to others (e.g., *map3k14* is inversely correlated to miR-7a-5p and miR-138-5p and directly correlated to miR-212-5p, while *bdnf* is inversely correlated to let-7b-3p and directly correlated to miR-212-5p). This observation prompts the hypothesis that some mRNAs may be subject to a dual control by different miRNAs at cytosolic and nuclear level. This hypothesis should be challenged and investigated.

Genes that are anticorrelated with miRNAs are enriched for "epileptogenic" ontology categories

To further investigate the functional role of miRNAs significantly differentially expressed and anticorrelated

with their predicted mRNA targets, we examined the functional enrichment of the mRNA targets identified in epileptogenesis and chronic phases of epilepsy.

Target genes that inversely correlated with differentially expressed miRNAs during the epileptogenesis period were enriched for GO terms related to synaptic function [like "response to stimulus" (p = 0.0013), "signaling" (p =2.68⁻⁰⁵), "signal transduction" (p = 0.0047), and others] and immunity [like "humoral immune response" (p =0.0009), "regulation of immune system process" (p = 0.0013), and others]. In addition, terms related to complement activation [like "complement activation" (p = 0.0002) and "complement activation, classical pathway" (p = 0.0009)] are in prominent position (Fig. 5A; Table 8). Proteins of the classical complement pathway not only play a role in the innate immune system, but have been also shown to be released from neurons, and serve as a new class of synaptic organizers (Yuzaki, 2017). These GO terms are potentially relevant to changes occurring at the level of the DG in epileptogenesis (Dudek and Sutula, 2007; Vezzani

Table 6. miRNA-mRNA fold changes positive correlation in epileptogenesis

	microRNA data				mRNA targets of respective microRNA (FDR $<$ 0.1)								
miRNA	Guiù	Perforant path	Amygdala					Perfora path	ant	Amygd	ala	Kainic	
Name	Pilocarpine	stimulation	stimulation		mRNA gene names	Pilocar	oine	stimula	ition	stimula		acid	
				Meta-analysis		50	Adjusted	50	Adjusted	50	Adjusted	50	Adjuste
miD 202 En	-0.50	miRNA FC -0.35	-0.75	<i>p</i> value 6.11 ⁻⁰⁵	Road?	FC -0.58	p value 0.0038	FC -1.24	<i>p</i> value 0.0012	FC -0.59	<i>p</i> value 0.0010	FC -0.87	<i>p</i> value 0.0034
miR-383-5p	-0.50	-0.35	-0.75	0.11	Rasd2 Sec14l1	-0.58	0.0038	-0.50	0.0012	-0.39	0.0010	-0.87	0.0034
					Мрр6	-0.59	0.0295	-0.54	0.0517	-0.39	0.4123	-0.78	0.0510
miR-153-3p	-0.32	-0.42	-0.22	0.0008	Mthfd1I	-1.27	0.0398	-0.83	0.0102	-0.94	0.0005	-0.44	0.0093
					Gdf10	-2.91	8.62^{-06}	-1.46	0.0306	-1.86	2.27^{-07}	-1.02	0.0014
					Nr4a3	0.12	0.9726	-1.65	0.0721	-1.64	0.0574	-0.65	0.0042
					Mettl7a	-0.74	0.0066	-0.72	0.0861	-0.87	0.0750	-0.19	0.0500
	0.00	0.00	0.00	0.550-06	Ablim2	-0.05	0.9953	-0.78	0.0964	-0.35	0.7323	-0.29	0.0667
miR-324-5p miR-150-5p	-0.09 -0.45	-0.60 -0.45	-0.29 -0.56	8.552 ⁻⁰⁶ 6.168 ⁻⁰⁵	Ryr1 Ddit4l	-1.52 -2.38	0.0022 0.0063	-1.39 -1.89	0.0196 0.0098	-1.86 -2.00	2.53 ⁻⁰⁸ 6.09 ⁻⁰⁶	-0.68 -1.09	0.0014 0.0007
min-150-5p	-0.45	-0.45	-0.50	0.100	Htr5b	-1.55	3.87 ⁻⁰⁵	-1.66	0.0263	-0.54	0.0096	-0.73	0.0016
					Fkbp4	-0.21	0.7718	-0.55	0.0464	-0.45	0.0737	-0.26	0.0775
					Pip5k1b	-1.03	0.0228	-0.46	0.0492	-0.79	0.1731	-0.38	0.0619
					Calml4	-0.60	0.3155	-0.44	0.0839	-0.48	0.1135	-0.38	0.0139
miR-92b-3p	-1.05	-0.72	-0.37	1.695^{-05}	Per2	-0.59	0.8691	-0.46	0.0717	-0.56	0.0253	-0.37	0.0918
miR-345-5p	-0.23	-0.25	-0.12	1.464^{-06}	Rasd2	-0.58	0.0038	-1.24	0.0012	-0.59	0.0010	-0.87	0.0034
					Ryr1	-1.52	0.0022	-1.39	0.0196	-1.86	2.53 ⁻⁰⁸	-0.68	0.0014
					Kihi14	-2.36	2.39 ⁻⁰⁶	-1.66	0.0204	-0.84	0.0006	-0.44	0.0077
					B3galt5 Pip5k1b	-1.93 -1.03	0.0118 0.0228	-0.83 -0.46	0.0442 0.0492	-1.39 -0.79	7.49 ⁻⁰⁵ 0.1731	-0.43 -0.38	0.0471 0.0619
					Rspo3	-0.41	0.0228	-0.40	0.0492	-0.79	0.1731	-0.55	0.0259
					Fat1	-1.55	0.0012	-1.01	0.0020	-1.05	0.0012	-0.19	0.0792
miR-101a-3p	-0.68	-0.71	-0.19	5.148 ⁻⁰⁶	Rasd2	-0.58	0.0038	-1.24	0.0012	-0.59	0.0010	-0.87	0.0034
					Ddit4l	-2.38	0.0063	-1.89	0.0098	-2.00	6.09^{-06}	-1.09	0.0007
					Gdf10	-2.91	8.62^{-06}	-1.46	0.0306	-1.86	2.27^{-07}	-1.02	0.0014
					Plk5	-2.15	0.0033	-0.59	0.0311	-1.46	0.0013	-1.00	0.0303
					Plag1	-0.58	0.0528	-0.45	0.0425	0.06	0.5075	-0.59	0.0453
miR-29c-5p	-0.59	-0.41	-0.14	1.477^{-07}	Crim1	-1.07	0.0084	-0.59	0.0173	-0.87	9.88 ⁻⁰⁶	-0.31	0.0323
					Dnah12 C5ar1	-0.35 0.43	0.9054 0.2817	0.65 0.33	0.0492 0.3103	0.32 0.44	0.4073	0.22 0.24	0.0340 0.0841
					Slc20a1	0.43	0.2817	0.33	0.5103	0.44	0.0181 0.0227	0.24	0.0041
miR-330-3p	-0.99	-0.40	-0.32	1.935 ⁻⁰⁶	Gabrd	-0.71	0.0282	-0.59	0.0067	-0.53	0.0053	-0.55	0.0070
					Ets2	-0.44	0.3991	-0.66	0.0249	-0.44	0.0641	-0.31	0.0095
					Gpc3	-1.92	0.0026	-0.85	0.0492	-0.89	0.0006	-1.29	0.0054
miR-138-5p	-0.48	-0.29	-0.50	3.983^{-08}	Rasd2	-0.58	0.0038	-1.24	0.0012	-0.59	0.0010	-0.87	0.0034
					Nr4a1	0.24	0.7205	-0.88	0.0125	-0.56	0.4123	-0.32	0.0719
					Crim1	-1.07	0.0084	-0.59	0.0173	-0.87	9.88^{-06}	-0.31	0.0323
					Nhlh1	-1.67	0.0023	-1.12	0.0337	-0.63	0.0295	-0.84	0.0097
	0.04	0.00	0.40	1.62^{-07}	Nr4a3	0.12	0.9726	-1.65	0.0721 0.0012	-1.64	0.0574	-0.65	0.0042
miR-667-3p	-0.34	-0.20	-0.49	1.02	Rasd2 Etv5	-0.58 -0.08	0.0038 0.9646	-1.24 -0.63	0.0012	-0.59 -0.47	0.0010 0.2965	-0.87 -0.49	0.0034 0.0044
miR-212-5p	2.55	0.27	0.28	1.152 ⁻⁰⁵	Sox11	1.52	0.0027	1.36	0.0092	1.57	0.2303	1.23	0.0011
					Serping1	1.15	0.1500	1.22	0.0204	1.19	0.0034	2.12	0.0009
					Map3k14	0.20	0.8682	0.77	0.0263	0.11	0.0834	0.26	0.0653
					Ptprn	0.83	0.0001	0.56	0.0613	0.38	0.0034	0.33	0.0648
					Kank2	0.00	0.9473	0.71	0.0673	0.62	0.0153	0.59	0.0241
					Acan	0.30	0.0723	0.37	0.0799	0.17	0.2577	0.37	0.0436
					SIc7a14	0.35	0.1717	0.42	0.0877	0.91	0.0003	0.23	0.0568
					C1qc	0.92 0.58	0.1759 0.0494	0.42 0.40	0.3397 0.4124	1.11 0.87	0.0014 0.0299	0.97 0.97	0.0816 0.0299
					Ly86 Slc20a1	0.58	0.0494	0.40	0.4124	0.87	0.0299	0.97	0.0299
					Blnk	0.27	0.0039	0.43	0.1925	0.71	0.0227	0.43	0.0042
					Bdnf	1.17	0.0297	0.20	0.6933	0.45	0.0352	0.32	0.0487
					Pdlim4	0.19	0.7034	0.15	0.5799	0.15	0.0771	0.46	0.0721
			Syn2	0.39	0.0258	0.55	0.0299	0.65	0.0399	0.11	0.2134		
					Epb41l4b	0.57	0.0030	0.58	0.0237	0.70	0.0008	0.17	0.4555
					Nnat	0.63	0.2970	0.86	0.0717	1.09	0.0083	0.15	0.5894
					Htatip2	0.79	0.0007	0.37	0.0669	0.70	0.0119	-0.02	0.9557
					Trh	2.87	0.0096	1.31	0.0246	1.20	0.0145	0.99	0.3361
lot_7b_2s	-0.80	-0.41	5 75	8.364 ⁻⁰⁶	Asph Pspo3	0.05 -0.41	0.9026 0.5671	0.45 -0.41	0.0254 0.0626	0.45 -0.11	0.0832	0.17 -0.55	0.1308
let-7b-3p	-0.00	-0.41	-5.75	0.004	Rspo3	-0.41	0.3071	-0.41	0.0020	-0.11	0.1711	-0.05	0.0259

Table 6. Continued

	microRNA data				mRNA targets microRNA (FE								
miRNA Name	Pilocarpine	Perforant path stimulation	Amygdala stimulation		mRNA gene names	Pilocar	nino	Perfora path stimula		Amygd stimula		Kainic acid	
Name	Filocarpine	Stimulation	Sumulation	Meta-analysis	Tiames	Filocal	Adjusted	Sumula	Adjusted	Sumula	Adjusted	aciu	Adjuste
		miRNA FC		p value		FC	p value	FC	p value	FC	p value	FC	p value
miR-132-3p	1.91	0.80	0.10	4.195 ⁻¹⁸	Efr3a	1.11	4.17 ⁻⁰⁷	1.40	0.0005	1.28	0.0096	0.49	0.0070
1111 102 Op	1.01	0.00	0.10	4.100	Sox11	1.52	0.0027	1.36	0.0092	1.57	0.0063	1.23	0.0011
					Wis	1.75	1.52 ⁻⁰⁵	0.80	0.0108	1.45	7.49 ⁻⁰⁵	0.57	0.0051
					Rin2	-0.06	0.9546	0.43	0.0517	0.57	0.0638	0.44	0.0303
					Gpnmb	0.81	0.2150	0.30	0.6840	1.54	0.0030	1.45	0.0259
					Zfp521	0.59	0.0031	0.74	0.0052	1.23	0.0002	0.23	0.2169
					Asph	0.05	0.9026	0.45	0.0254	0.45	0.0832	0.17	0.1308
miR-146a-5p	4.71	0.37	0.33	4.047 ⁻¹⁰	Mdm1	0.00	0.0680	0.76	0.0234	0.69	0.0360	0.30	0.0322
mine 140a-op	4.71	0.07	0.00	4.047	Inpp4b	0.23	0.1291	1.15	0.0027	0.80	0.1003	0.31	0.0735
					Arhgap17	0.33	0.0762	0.88	0.0027	0.30	0.0183	0.43	0.0259
					Igsf1	0.50	0.4934	0.89	0.0000	1.03	0.3719	0.43	0.0239
					Gpat3	0.30	0.4934	0.89	0.0423	-0.04	0.2317	0.16	0.0923
					Sowahc	0.44	0.4497	0.40	0.3070	-0.04 1.09	0.2317	0.10	0.0923
					Fcer1g	1.09	0.0003	0.45	0.1380	1.14	0.0008	1.37	0.0073
					Slfn13	0.40	0.0112	0.31	0.1380	0.46	0.0002	1.53	0.0039
						0.40	0.0031	0.41	0.0052	1.23	0.0429	0.23	0.2169
					Zfp521 Anks1a		0.5280	0.74		0.72	0.0002	0.23	0.2169
						0.17			0.0492				
					Trh	2.87	0.0096	1.31	0.0246	1.20	0.0145	0.99	0.3361
					Ntm	-0.32	0.9428	0.37	0.0861	0.71	0.0527	-0.22	0.3746
	0.70	0.05	0.40	0.000-14	Col9a1	1.28	0.0003	0.91	0.0425	0.41	0.0208	0.03	0.9192
miR-551b-3p	-0.78	-0.85	-0.42	9.092^{-14}	Plxdc1	-0.70	0.0007	-0.73	0.0127	-0.28	0.0555	-0.33	0.0823
					Ogfrl1	-0.23	0.1736	-0.52	0.0135	-0.55	0.0293	-0.30	0.0563
					Htr5b	-1.55	3.87 ⁻⁰⁵	-1.66	0.0263	-0.54	0.0096	-0.73	0.0016
	0.74	0.44		5 400-14	Diaph1	-0.27	0.2293	-0.30	0.0492	0.07	0.3960	-0.23	0.0530
miR-344b-2-3p	3.74	0.11	0.29	5.432 ⁻¹⁴	Runx1	1.22	0.0009	1.48	0.0027	1.08	0.0008	0.60	0.0027
					ll18	0.31	0.3608	0.52	0.0669	0.38	0.1009	0.51	0.0955
					Chi3l1	0.33	0.8228	0.02	0.9731	1.14	0.0881	0.31	0.0785
					Fancd2os	0.85	0.0043	0.36	0.1786	0.66	0.0353	0.28	0.0372
					P2ry6	0.50	0.1614	0.12	0.7169	0.17	0.0499	0.61	0.0231
				17	Asph	0.05	0.9026	0.45	0.0254	0.45	0.0832	0.17	0.1308
miR-139-5p	-1.40	-0.77	-0.91	4.4^{-17}	Gabrd	-0.71	0.0282	-0.59	0.0067	-0.53	0.0053	-0.55	0.0070
					Gdf10	-2.91	8.62^{-06}	-1.46	0.0306	-1.86	2.27^{-07}	-1.02	0.0014
				10	Rspo3	-0.41	0.5671	-0.41	0.0626	-0.11	0.1711	-0.55	0.0259
miR-33-5p	-2.40	-0.80	-0.51	4.916 ⁻¹⁹	Smarca2	-0.60	0.2192	-1.53	0.0186	-0.35	0.0904	-0.27	0.0340
					Fxyd7	-1.49	7.16^{-05}	-1.65	0.0186	-0.98	0.0052	-0.30	0.0142
					Arg1	-1.24	0.0324	-0.49	0.0984	-0.37	0.0679	-0.50	0.0344
miR-7a-5p	-0.91	-0.93	-0.51	3.436 ⁻²⁴	Hpca	-0.40	0.0880	-0.64	0.0090	-0.74	0.0220	-0.22	0.0244
miR-212-3p	1.09	0.88	0.22	1.744^{-47}	Efr3a	1.11	4.17^{-07}	1.40	0.0005	1.28	0.0096	0.49	0.0070
					Sox11	1.52	0.0027	1.36	0.0092	1.57	0.0063	1.23	0.0011
					Wls	1.75	1.52^{-05}	0.80	0.0108	1.45	7.49^{-05}	0.57	0.0051
					Rin2	-0.06	0.9546	0.43	0.0517	0.57	0.0638	0.44	0.0303
					Gpnmb	0.81	0.2150	0.30	0.6840	1.54	0.0030	1.45	0.0259
					Zfp521	0.59	0.0031	0.74	0.0052	1.23	0.0002	0.23	0.2169
					Asph	0.05	0.9026	0.45	0.0254	0.45	0.0832	0.17	0.1308

et al., 2015). In addition, at the level of cell signaling pathways, analysis of KEGG pathways enriched among the mRNA targets suggested a key role for the MAPK cascade (p = 0.354; Fig. 5B; Table 8). Notably, changes in the activation state of kinase pathways and altered kinase expression patterns have been reported in the hippocampus by previous studies (Xi et al., 2009). In the chronic period, the predicted and anticorrelated mRNA targets revealed enrichment in biological processes that have been previously implicated in chronic epilepsy (Ludewig et al., 2016; Robel and Sontheimer, 2016) such as "regulation of dendritic cell differentiation" ($p = 4.8 \times 10^{-6}$), "glial cell development" ($p = 4.0 \times 10^{-4}$), "proliferation" (p $= 6.0 \times 10^{-4}$), and "cell proliferation" ($p = 4.0 \times 10^{-4}$; Fig. 5C; Table 9). Notably, we performed a permutation test to check for false positive enrichment in GO term and KEGG pathway analysis, but this did not change any of the results.

Target genes that directly correlated with differentially expressed miRNAs during the epileptogenesis period were enriched for GO terms related to glia proliferation ["regulation of glial cell proliferation" (p = 0.0001) and "glial cell proliferation" (p = 0.0001)]. In addition, and as in the inverse correlation analysis, terms related to complement activation [like "complement activation, classical pathway" (p = 0.0004)] were significantly enriched. In the chronic period, the positively correlated mRNA targets revealed significant (p < 0.0004)] enrichment in GO terms related to receptor function like "receptor activity," "signaling receptor activity," "molecular transducer activity," and "transmembrane signaling receptor activity."

Table 7. miRNA-mRNA fold changes positive correlation in the chronic period

	microRNA data				mRNA targets of respective microRNA (FDR $<$ 0.1)			
miRNA name	Pilocarpine	Perforant path stimulation	Amygdala stimulation		mRNA gene names	Amygdala stimulation	Amygdala stimulatior	
	1 liocalpino	otimulation	otimolation	Meta-analysis	namoo	otimulation	Adjusted	
		FC		p value		FC	p value	
miR-652-3p	-0.90	-0.34	-0.41	9.65 ⁻⁰⁷	Ano2	-0.39	0.0077	
					Ece2	-0.22	0.0395	
					Optn	-0.26	0.0825	
niR-551b-3p	-1.30	-0.52	-0.50	0.0006	Clmp	-0.36	0.0325	
					Socs5	-0.36	0.0619	
					Asic2	-0.26	0.0731	
niR-324-3p	-0.44	-0.22	-0.27	0.0055	Gdf10	-0.94	0.0083	
					Gpr176	-0.43	0.0116	
					Etv5	-0.58	0.0202	
					Elfn2	-0.25	0.0225	
					Tcerg1l	-0.42	0.0311	
					Sstr3	-0.33	0.0371	
					Nr4a3	-0.50	0.0372	
					Veph1	-0.27	0.0380	
					Cyp26b1	-0.35	0.0380	
					Itgb4	-0.50	0.0394	
					Nefm	-0.31	0.0394	
					Alcam	-0.29	0.0547	
					Grik1	-0.30	0.0555	
					Clmn	-0.30	0.0619	
					Arg1	-0.30	0.0697	
					Grik3	-0.72	0.0702	
					Asic2	-0.26	0.0731	
					Boc	-0.33	0.0772	
					Ubash3b	-0.24	0.0772	
						-0.24	0.0817	
	0.04	-0.29	-0.41	0.0317	Cbarp Dis Eld	-0.50	0.0077	
niR-148b-3p	-0.24	-0.29	-0.41	0.0317	Pip5k1b	-0.43	0.0077	
					Gpr176 Slit2	-0.43 -0.39	0.0118	
					Camk1g	-0.43	0.0234	
					Gpr165	-0.48	0.0326	
					Hcrtr2	-0.28	0.0371	
					Htra4	-1.37	0.0380	
					Vstm2b	-0.40	0.0504	
					Alcam	-0.29	0.0547	
					Ankrd34c	-0.44	0.0568	
					Ppara	-0.25	0.0773	
ID (00 -			o (o	0.0407	Hnrnpm	-0.22	0.0988	
niR-130a-3p	-0.24	-0.32	-0.46	0.0107	Eloc	-0.33	0.0380	
					Htra4	-1.37	0.0380	
					Trhr	-0.75	0.0380	
					Mthfd11	-0.39	0.0505	
					Rasd2	-0.47	0.0560	

To infer the functional relationships between miRNAs identified as differentially expressed in the meta-analysis, we created a network of miRNAs based on their predicted anticorrelated target pathways (Fig. 5). These results highlight that several distinct miRNAs may contribute to the regulation of functionally related processes and pathways, and so prioritizing individual miRNAs as potential therapeutic targets will require downstream experimental analysis.

Discussion

Main findings

The present meta-analysis provides a miRNA differential expression signature in the DG of rats during epileptogenesis and in the chronic phase of epilepsy. We identified 26 miRNAs significantly differentially expressed during epileptogenesis, and five miRNAs significantly differentially expressed in the chronic phase of epilepsy. We also identified 11 miRNAs in epileptogenesis and two in chronic epilepsy that were identified as significantly differentially expressed by the meta-analysis but not in any of the individual studies. Further, we explored the negative correlation between the significantly differentially expressed miRNAs and their predicted mRNA targets in the same models of epilepsy. We identified 122 predicted mRNAs targets with an anticorrelated expression relationship to 22 of the 26 miRNAs significantly differentially

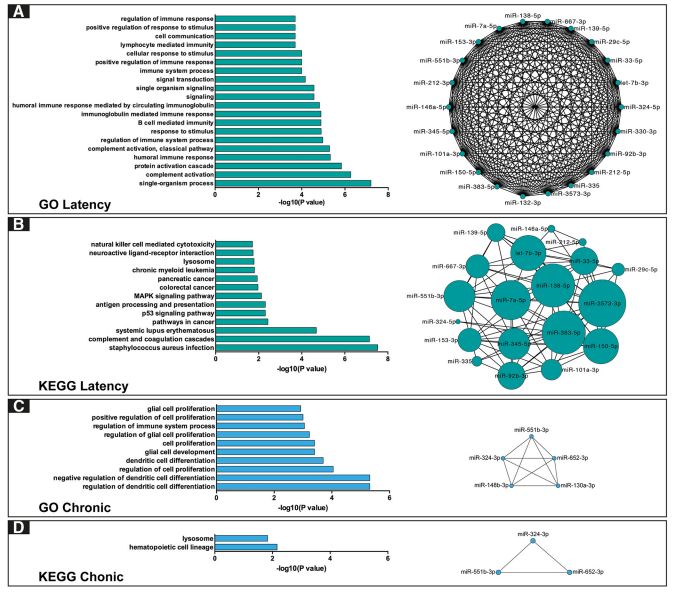


Figure 5. Functional enrichment of dysregulated miRNA-mRNAs targets modules. *A*, Horizontal bar plots (on the left) show the GO enrichment status (top 20 terms) for 112 predicted mRNAs targets that anticorrelate with 22 miRNAs expression level in epileptogenesis (FDR < 5%, hypergeometric test). The miRNA-mRNA module is represented by a network graph (on the right) showing the connections between miRNAs based on the function of their mRNAs predictive targets revealed by the GO enrichment. *B*, Horizontal bar plots (on the left) show KEGG enrichment analysis for predicted mRNAs targets that anticorrelate with miRNAs expression level in epileptogenesis (FDR < 5%, hypergeometric test). miRNA-mRNA modules are represented with network plot (on the right) showing the connection between miRNAs based on the pathways in which are involved their predicted targets revealed by KEGG analysis. *C*, *D*, GO and KEGG enrichment status for 29 predicted miRNA targets that anticorrelate with five miRNAs differentially expressed in the chronic stage (FDR < 5%, hypergeometric test).

expressed in epileptogenesis. Below, we discuss these findings and their possible implications in the development and maintenance of epilepsy. Together, we also discuss the intrinsic limitations of this study that must be taken into account.

Epileptogenesis

Functional annotations of the target genes of miRNAs significantly differentially expressed during the latent interval between brain injury and the development of spontaneous seizures (epilepsy) support a relationship between dysregulated miRNAs and molecular and cellular reorganizations that are known to occur during epileptogenesis. First, the GO enrichment analysis identifies many terms that suggest a role for modulation of synaptic transmission during epileptogenesis. This is not surprising, given the critical role of the DG in the temporal lobe seizure network (Krook-Magnuson et al., 2015) and previous experimental evidence for changes in synaptic efficacy and connections during epileptogenesis (Dudek and Sutula, 2007). Another set of terms broadly refers to immunity and inflammation, events that are deeply asso-

Table 8. GO and KEGG enrichment of 122 predicted and anticorrelated mRNAs targets of 22 miRNAs differentially expressed in epileptogenesis

Biological process $Bingle-graniem = 0000449 = 0 - 2499 = 0 - 67 = 1 - 0.211 = 1 - 1.29 = 1.20 + 0.21 = 0.0000000000000000000000000000000000$	Biological processSingle-organism processGO:004469C = 2489O = 67E = 42.21R = 1.59Biological processComplement activationGO:000695C = 11O = 5E = 0.19R = 26.80Biological processProtein activationGO:007237C = 13O = 5E = 0.22R = 22.68Biological processProtein activationGO:000695C = 16O = 5E = 0.27R = 18.43Biological processHumoral immuneGO:000695C = 8O = 4E = 0.14R = 29.48Biological processComplement activation, classical pathwayGO:001972C = 32O = 6E = 0.54R = 11.06Biological processRegulation of immune system processGO:00268C = 242O = 15E = 4.10R = 3.65Biological processResponse to stimulusGO:005089C = 2294O = 59E = 38.90R = 1.52	6.29^{-08} 5.34^{-07} 1.45^{-06} 4.72^{-06} 5.14^{-06}	0.0004 0.0009
boxes boxes Biological process Protes activation GOU00005 C = 11 O = 5 E = 0.19 R = 28.80 1.45^{-100} 0.0 Biological process Protes activation GOU000055 C = 16 O = 5 E = 0.127 R = 12.48 4.12 ⁻¹²⁰ 0.0 Biological process Beal resultation GOU000055 C = 45 O = 4 E = 0.14 R = 28.48 5.14 ⁻¹²⁰ 0.0 Biological process Beal resultation GOU000055 C = 42 O = 4 E = 0.14 R = 11.60 1.28 ⁻¹²⁰ 0.0 Biological process Regulation of minute GOU000050 C = 224 O = 4 E = 0.54 R = 11.60 1.28 ⁻¹²⁰ 0.0 Biological process Regulation of minute segments GOU000050 C = 2320 O = 4 E = 0.54 R = 11.26 1.28 ⁻¹²⁰ 0.0 Biological process Regulation of minute segments GOU00010 C = 1010 O = 4 E = 0.54 R = 11.26 1.28 ⁻¹²⁰ 0.0 Biological process Signafing ments	processBiological processComplement activationGO:000695C = 11O = 5E = 0.19R = 26.80Biological processProtein activationGO:007237C = 13O = 5E = 0.22R = 22.68Biological processHumoral immuneGO:000695C = 16O = 5E = 0.27R = 18.43Biological processComplement activation, classical pathwayGO:000695C = 8O = 4E = 0.14R = 29.48Biological processB cell-mediated immunityGO:001972C = 32O = 6E = 0.54R = 11.06Biological processRegulation of immune system processGO:00268C = 242O = 15E = 4.10R = 3.65Biological processResponse to stimulusGO:005089C = 2294O = 59E = 38.90R = 1.52	5.34 ⁻⁰⁷ 1.45 ⁻⁰⁶ 4.72 ⁻⁰⁶ 5.14 ⁻⁰⁶	0.0002 0.0004 0.0009 0.0009
Biological process Complement advantom GOUDDES C = 11 O = 5 E = 0.19 R = 28.80 1.45^{-50} D.00 Biological process Proces advance Complexes D = 5 E = 0.27 R = 18.40 4.22^{-56} D.0 Biological process D = 5 E = 0.27 R = 18.40 1.28^{-50} D.0 Biological process B estimation of immune GOUDD028 C = 4 O = 4 E = 0.54 R = 11.00 1.28^{-50} D.0 Biological process Requiring of immune GOUDD028 C = 220 D = 6 E = 0.54 R = 11.00 1.28^{-50} D.0 Biological process Requiring of immune method GOUDD028 C = 220 D = 6 E = 0.54 R = 11.00 1.28^{-50} D.0 Biological process Requiring of immune method GOUDD025 C = 10 D = 4 E = 0.51 R = 17.2 2.68^{-50} D.0 Biological process Signaling one GOUDD027 C = 108 D = 4 E = 0.51 R = 1.72 2.68^{-50} D.0 D =	Biological processComplement activationGO:000695C = 11O = 5E = 0.19R = 26.80Biological processProtein activationGO:007237C = 13O = 5E = 0.22R = 22.68Biological processHumoral immuneGO:000695C = 16O = 5E = 0.27R = 18.43Biological processComplement activation, classical pathwayGO:000695C = 8O = 4E = 0.14R = 29.48Biological processB cell-mediated immunityGO:001972C = 32O = 6E = 0.54R = 11.06Biological processRegulation of immune system processGO:00268C = 242O = 15E = 4.10R = 3.65Biological processResponse to stimulusGO:005089C = 2294O = 59E = 38.90R = 1.52	1.45^{-06} 4.72^{-06} 5.14^{-06}	0.0004 0.0009
Biological process Protein astivation CO.007237 C = 13 O = 5 E = 0.22 R = 2.2.8 1.45 Co. Biological process Humonal Immune 60000000 C = 16 O = 5 E = 0.27 R = 16.43 A 12 ⁻²⁰ 0.0 Biological process Complement astivation, CO.0000000 C = 242 O = 6 E = 0.14 R = 2.8.40 1.45 ⁻⁴⁰ 0.0 Biological process Regonator of immune CO.0002000 C = 242 O = 6 E = 0.14 R = 2.8.40 1.50 ⁻⁴⁰ 0.0 Biological process Resconte to simulue CO.0002000 C = 242 O = 6 E = 0.14 R = 1.162 1.25 ⁻⁴⁰ 0.0 Biological process Resconte to simulue CO.0002000 C = 100 O = 4 E = 0.54 R = 1.72 2.64 ⁻⁴⁰ 0.0 Biological process Synghe-magnetim CO.000200 C = 110 O = 4 E = 2.510 R = 1.72 2.64 ⁻⁴⁰ 0.0 Biological process Synghe process Synghe process CO.000077 C = 150 O =	Biological processProtein activation cascadeGO:007237C = 13O = 5E = 0.22R = 22.68Biological processHumoral immune responseGO:000695C = 16O = 5E = 0.27R = 18.43Biological processComplement activation, classical pathwayGO:000695C = 8O = 4E = 0.14R = 29.48Biological processComplement activation, classical pathwayGO:001972C = 32O = 6E = 0.54R = 11.06Biological processB cell-mediated immunityGO:00268C = 242O = 15E = 4.10R = 3.65Biological processResponse to stimulusGO:005089C = 2294O = 59E = 38.90R = 1.52	1.45^{-06} 4.72^{-06} 5.14^{-06}	0.0004
cascade constant	Biological processHumoral immune responseGO:000695C = 16O = 5E = 0.27R = 18.43Biological processComplement activation, classical pathwayGO:000695C = 8O = 4E = 0.14R = 29.48Biological processB cell-mediated immunityGO:001972C = 32O = 6E = 0.54R = 11.06Biological processRegulation of immune system processGO:00268C = 242O = 15E = 4.10R = 3.65Biological processResponse to stimulusGO:005089C = 2294O = 59E = 38.90R = 1.52	4.72 ⁻⁰⁶ 5.14 ⁻⁰⁶	0.0009
Humon immune GO.00065 C = 10 O = 5 E = 0.27 R = 16.43 4.72 ⁻⁶⁵ 0.0 Biological process Complement activation, obstacial proteins GO.000585 C = 8 O = 4 E = 0.14 R = 29.48 5.14 ⁻⁶⁵ 0.0 Biological process B cell mediated GO.000585 C = 242 O = 15 E = 0.14 R = 29.48 5.14 ⁻⁶⁵ 0.0 Biological process Pergenomes CO.000580 C = 2240 O = 15 E = 0.14 R = 10.05 1.25 ⁻⁶⁵ 0.0 Biological process Pergenomes/and immune response CO.000245 C = 10 O = 4 E = 0.17 R = 1.72 2.85 ⁻⁶⁵ 0.0 Biological process Signaling CO.002245 C = 1510 O = 44 E = 28.61 R = 1.72 2.85 ⁻⁶⁵ 0.00 Biological process Signaling CO.002245 C = 1510 O = 44 E = 28.61 R = 1.72 2.85 ⁻⁶⁵ 0.00 Biological proces Signaling CO.0021470 C = 1510 O = 44 E = 28.61 R = 1.62	Biological processHumoral immune responseGO:000695C = 16O = 5E = 0.27R = 18.43Biological processComplement activation, classical pathwayGO:000695C = 8O = 4E = 0.14R = 29.48Biological processB cell-mediated immunityGO:001972C = 32O = 6E = 0.54R = 11.06Biological processRegulation of immune system processGO:00268C = 242O = 15E = 4.10R = 3.65Biological processResponse to stimulusGO:005089C = 2294O = 59E = 38.90R = 1.52	5.14 ⁻⁰⁶	
Integrate <	response Biological process Complement activation, classical pathway Biological process B cell-mediated G0:001972 C = 32 O = 6 E = 0.54 R = 11.06 immunity Biological process Regulation of immune G0:000268 C = 242 O = 15 E = 4.10 R = 3.65 system process Biological process Response to stimulus G0:005089 C = 2294 O = 59 E = 38.90 R = 1.52	5.14 ⁻⁰⁶	
Biological process Complement activation, classical pathway COUNDERS C = 8 C = 4 E = 0.14 R = 28.48 S.14 ⁻⁵⁰ C 0.200000000000000000000000000000000000	Biological processComplement activation, classical pathwayGO:000695C = 8O = 4E = 0.14R = 29.48Biological processB cell-mediated immunityGO:001972C = 32O = 6E = 0.54R = 11.06Biological processRegulation of immune system processGO:000268C = 242O = 15E = 4.10R = 3.65Biological processResponse to stimulusGO:005089C = 2294O = 59E = 38.90R = 1.52		0.0009
a classical pathway	classical pathway Biological process B cell-mediated GO:001972 C = 32 O = 6 E = 0.54 R = 11.06 immunity Biological process Regulation of immune system process GO:000268 C = 242 O = 15 E = 4.10 R = 3.65 Biological process Response to stimulus GO:005089 C = 2294 O = 59 E = 38.90 R = 1.52		
Biological process B cell-modified G 0,001972 C - 32 O - 6 E - 0.64 R - 11.00 1.28 ⁻⁴⁰ 0.0 Biological process Pepter process O - 15 E = 4.10 R = 3.85 1.55 ⁻⁵⁰ 0.0 Biological process Pepter process O - 58 E - 38.30 R = 1.52 1.25 ⁻⁵⁰ 0.0 Biological process Immuno response G 0.000245 C = 10 O = 4 E = 0.51 R = 1.72 2.86 ⁻⁵⁰ 0.0 Biological process Single-organism G 0.000245 C = 110 O = 44 E = 25.61 R = 1.72 2.86 ⁻⁵⁰ 0.0 Biological process Single-organism G 0.000716 C - 1309 O - 24 E = 25.61 R = 1.72 2.86 ⁻⁵⁰ 0.00 Biological process Single-organism G 0.000716 C - 1309 O - 19 E - 23.68 R - 1.76 6.75 ⁻⁵⁰ 0.00 Biological process Caluer response G 0.000717 C - 1711 O - 46 E - 29.02 R - 1.59 0.0001 0.0 Biological process Caluer response G 0.000777 C - 100 O - 8 <td< td=""><td>Biological process B cell-mediated immunity GO:001972 C = 32 O = 6 E = 0.54 R = 11.06 Biological process Regulation of immune system process GO:000268 C = 242 O = 15 E = 4.10 R = 3.65 Biological process Response to stimulus GO:005089 C = 2294 O = 59 E = 38.90 R = 1.52</td><td>1.28⁻⁰⁵</td><td></td></td<>	Biological process B cell-mediated immunity GO:001972 C = 32 O = 6 E = 0.54 R = 11.06 Biological process Regulation of immune system process GO:000268 C = 242 O = 15 E = 4.10 R = 3.65 Biological process Response to stimulus GO:005089 C = 2294 O = 59 E = 38.90 R = 1.52	1.28 ⁻⁰⁵	
Immunity Biological process Regulation of minume S0 000289 C = 242 O = 15 E = 4.10 R = 3.85 1.05 0.0 Biological process Immunolopulin-mediated GO 000100 C = 32 O = 6 E = 0.54 R = 1.12 1.25 0.0 Biological process Humoral immure response GO 000246 C = 10 O = 4 E = 0.17 R = 23.99 1.00 75 0.0 Biological process Single-origonic GO 0002470 C = 1510 O = 44 E = 26.61 R = 1.72 2.88 63 0.0 Biological process Single-origonic GO 000470 C = 1510 O = 44 E = 26.61 R = 1.72 2.88 63 0.00 Biological process Signal anduction GO 000277 C = 446 O = 39 E = 2.18 R = 1.72 0.0001 0.0 Biological process Immune system process GO 0002715 C = 1570 O = 45 E = 2.02 R = 1.52 0.0002 0.0 Biological process C colosass	immunity Biological process Regulation of immune GO:000268 C = 242 O = 15 E = 4.10 R = 3.65 system process Biological process Response to stimulus GO:005089 C = 2294 O = 59 E = 38.90 R = 1.52		0.0013
Biological process Perspectation of immune CO.00288 C = 242 O = 15 E = 4.10 R = 3.85 1.65 ⁺⁰⁵ 0.0 Biological process Response to stimulus GO.00008 C = 2244 O = 68 E = 0.54 R = 1.52 1.55 ⁻⁹⁰ 0.0 Biological process Mumani Presponse to stimulus GO.00025 C = 10 O = 4 E = 0.51 R = 1.52 1.50 ⁻⁹⁰ 0.0 Biological process Signific and process Signific and process GO.00275 C = 1510 O = 44 E = 26.61 R = 1.72 2.68 ⁻⁹⁰ 0.0 Biological process Signific and process Signific and process GO.00277 C = 446 O = 139 E = 27.68 R = 1.72 2.68 ⁻⁹⁰ 0.0 Biological process Gignific and process GO.00277 C = 446 O = 19 E = 27.68 R = 1.52 0.0001 0.0 Biological process Calidar response GO.00277 C = 117 O = 6 E = 0.88 R = 6.34 0.0002 0.0 Biological process Calidar response	Biological process Regulation of immune system process GO:000268 C = 242 O = 15 E = 4.10 R = 3.65 Biological process Response to stimulus GO:005089 C = 2294 O = 59 E = 38.90 R = 1.52		
a system process 1.25 ⁻⁰⁵ 1.25 ⁻⁰⁵ 0.0000000 0.0000000 0.0000000 0.00000000 0.0000000000000000000000000000000000	system process Biological process Response to stimulus GO:005089 C = 2294 O = 59 E = 38.90 R = 1.52	1.05^{-05}	0.0013
Biological process Persons to strutule GO.000289 C -2294 $O - 59$ E -8.30 R -1.52 1.25^{-35} 0.00 Biological process Immune response GO.000246 C -10 O -4 E -0.54 R -1.52 1.25^{-35} 0.00 multiple process Biological process GO.000245 C -10 O -4 E -0.17 R -23.59 1.50^{-16} 0.00 multiple process Signaling GO.000246 C -1510 O -44 E -25.61 R -1.72 2.68^{-35} 0.00 Biological process Signaling GO.000247 C -1510 O -44 E -25.61 R -1.72 2.68^{-35} 0.00 Biological process Signaling GO.000277 C -190 O -39 E -22.18 R -1.72 2.68^{-35} 0.00 Biological process Signaling GO.000277 C -90 O -6 E -28.02 R -1.59 0.0001 0.00 Biological process Immune response GO.000237 C -146 O -19 E -7.56 R -2.51 0.0001 0.00 murule response GO.000237 C -1711 O -46 E -28.03 R -1.59 0.0001 0.00 murule response GO.000277 C $=90$ O -6 E -0.86 R -0.94 0.0002 0.00 murule response GO.000271 C -1711 O -46 E -28.03 R -1.82 0.0002 0.00 murule for the response GO.000271 C -1570 O -43 E $-2.66.3$ R -1.82 0.0002 0.00 biological process Gel communication GO.000715 C -1570 O -43 E $-2.66.3$ R -1.82 0.0002 0.00 biological process Gel communication GO.000717 C -125 O -9 E -2.12 R -4.25 0.0002 0.00 biological process Immune response GO.00028 C -120 O -9 E -2.10 R -4.42 0.0002 0.00 biological process Immune response GO.00028 C -120 O -9 E -2.10 R -4.42 0.0002 0.00 biological process Immune response GO.00028 C -120 O -9 E -2.10 R -4.42 0.0002 0.00 biological process Immune response GO.00028 C -100 O -3 E -0.15 R -18.20 0.0004 0.00 c an anomatic recontania GO.00024 C -9 O -3 E -0.15 R -18.20 0.0005 0.00 Biological process Immune response GO.000290 C -3 O -9 E -2.10 R -4.42 0.0002 0.00 c anomatic recontania GO.000240 C -9 O -3 E -0.15 R -18.20 0.0005 0.00 Biological process Immune response GO.000290 C -3 O -2 E -0.05 R -30.31 0.0008 0.00 GO.0000 anomatic recontania GO.000290 C -3 O -2 E $-$	Biological process Response to stimulus GO:005089 C = 2294 O = 59 E = 38.90 R = 1.52		
Biological process Immunoglobulin-mediated momune response GO 000245 C $= 0$ E $= 0.54$ R $= 11.06$ 1.28^{-10} $= 0.54$ Biological process Humoral immuno response GO 000245 C -10 0 -4 E -0.17 R -23.59 1.50^{-16} 0.0 Biological process Signified regramm GO 000246 C -1510 0 -44 E $=2561$ R -1.72 2.66^{-65} 0.0 Biological process Signified regramm GO 0002470 C $=1510$ 0 $=44$ E $=22.51$ R $=1.72$ 2.66^{-65} 0.0 Biological process Immune regrams process GO 0005177 C $=90$ R E $=2.38$ R $=0.00001$ 0.0 Biological process Lymphocyte mediated GO 000247 C $=170$ E E 8.64 0.0002 0.0 Biological process Colobiological pr		1.25^{-05}	0.0013
Immune response Biological process Humoel immune response GO.000245 C = 10 O = 4 E = 0.17 R = 23.59 1.50 ⁻⁹⁵ 0.00 Biological process Single-signaling GO.002405 C = 1510 O = 44 E = 25.61 R = 1.72 2.66 ⁻⁹⁰ 0.0 Biological process Single-signaling GO.00237 C = 446 O = 19 E = 7.85 R = 1.72 2.66 ⁻⁹⁰ 0.0 Biological process Synal transaction GO.000237 C = 446 O = 19 E = 7.83 R = 5.24 0.0001 0.0 Biological process Immune response GO.000277 C = 910 O = 6 E = 1.33 R = 5.94 0.0002 0.0 Biological process Calutor response GO.000244 C = 51 O = 6 E = 0.86 R = 1.62 0.0002 0.0 Biological process Calutor munication GO.000278 C = 1710 O = 43 E - 26.63 R = 1.62 0.0002 0.0 Biological process Immune response GO.000278 C = 120 <td></td> <td></td> <td>0.0013</td>			0.0013
mediated by circulating immunopolation mediated by circulating granting GC/002305 C = 1510 O = 44 E = 25.61 R = 1.72 2.86^{-90} 0.00 Biological process Signal Immandeviton GC/002305 C = 1510 O = 44 E = 25.61 R = 1.72 2.86^{-90} 0.00 Biological process Signal Immandeviton GC/002307 C = 1308 O = 39 E = 25.61 R = 1.72 2.86^{-90} 0.00 Biological process Signal Immande system process GC/002377 C = 406 O = 6 E = 25.61 R = 1.53 R = 5.24 0.0001 0.00 Biological process Cell carresponts GC/002715 C = 1771 O = 6 E = 0.68 R = 6.52 0.0002 0.00 Biological process Coll comminication GC/002715 C = 1570 O = 4 E = 2.643 R = 1.52 0.0002 0.00 Biological process Regulation of Immune GC/002707 C = 125 O = 9 E = 2.12 R = 4.25 0.0002 0.00 Biological process Regulation of Immune			
$ \begin{array}{c} \text{Immunopletulin} \\ \text{Biological process} & Signaling & GO.002205 & C = 1510 & O = 44 & E = 25.81 & R = 1.72 & 2.88^{-95} & 0.0 \\ \text{Biological process} & Signaling & GO.00470 & C = 1510 & O = 44 & E = 25.81 & R = 1.72 & 2.89^{-95} & 0.0 \\ \text{Biological process} & Signalinansduction & GO.000716 & C = 1038 & O = 39 & E = 2.18 & R = 1.74 & 6.75^{-95} & 0.0 \\ \text{Biological process} & Desitive regulation of & GO.000271 & C = 46 & O = 19 & E = 7.58 & R = 5.24 & 0.0001 & 0.0 \\ \text{Immune response } & GO.000227 & C = 46 & O = 19 & E = 7.58 & R = 5.24 & 0.0001 & 0.0 \\ \text{Immuno response } & GO.000244 & C = 51 & O = 6 & E = 0.88 & R = 6.94 & 0.0002 & 0.0 \\ \text{Introduction of stimulus} & GO.0002715 & C = 1570 & O = 43 & E = 26.83 & R = 1.62 & 0.0002 & 0.0 \\ \text{Biological process} & Cell communication & GO.000715 & C = 1570 & O = 43 & E = 26.83 & R = 1.62 & 0.0002 & 0.0 \\ \text{Biological process} & Cell communication & GO.000277 & C = 125 & O = 9 & E = 2.12 & R = 4.25 & 0.0002 & 0.0 \\ \text{Biological process} & Positive regulation of immune & GO.00027 & C = 120 & O = 9 & E = 2.12 & R = 4.24 & 0.0002 & 0.0 \\ \text{Biological process} & Regulation of immune & GO.00027 & C = 12 & O = 9 & E = 2.12 & R = 4.24 & 0.0002 & 0.0 \\ \text{Biological process} & Immune response & GO.000027 & C = 10 & O = 9 & E = 2.12 & R = 4.24 & 0.0002 & 0.0 \\ \text{Biological process} & Immune response & GO.000027 & C = 10 & O = 3 & E = 0.17 & R = 17.69 & 0.0005 & 0.0 \\ \text{Biological process} & Immune response based & GO.00027 & C = 9 & O = 3 & E = 0.17 & R = 17.69 & 0.0005 & 0.0 \\ \text{Biological process} & Response based & GO.00027 & C = 10 & O = 3 & E = 0.17 & R = 17.69 & 0.0005 & 0.0 \\ \text{Biological process} & Response based & GO.000270 & C = 3 & O = 2 & E = 0.05 & R = 38.31 & 0.0006 & 0.0 \\ \text{Biological process} & Response bolid & GO.000290 & C = 3 & O = 2 & E = 0.05 & R = 38.31 & 0.0006 & 0.0 \\ \text{Biological process} & Response bolid & GO.000290 & C = 3 & O = 2 & E = 0.05 & R = 38.31 & 0.0006 & 0.0 \\ \text{Biological process} & Response bolid & GO.000290 & C = 3 & O = 2 &$	Biological process Humoral immune response $GO:000245$ $C = 10$ $O = 4$ $E = 0.17$ $R = 23.59$	1.50 ⁻⁰⁵	0.0014
Biological process Signaling GO:002305 C = 1510 O = 44 E = 28.61 R = 1.72 2.81 $^{-05}$ 0.00 Biological process Signaling GO:00470 C = 1510 O = 44 E = 28.61 R = 1.72 2.81 $^{-05}$ 0.00 Biological process Signaling GO:00471 C = 1308 O = 39 E = 22.18 R = 1.76 6 75 $^{-05}$ 0.00 Biological process Immune system process GO:000257 C = 466 O = 19 E = 7.56 R = 2.51 0.0001 0.00 Immune response GO:000577 C = 90 O = 8 E = 1.53 R = 5.24 0.0001 0.00 Immune response GO:000214 C = 511 O = 6 E = 0.86 R = 6.94 0.0002 0.00 Biological process Immune response GO:000257 C = 150 O = 6 E = 0.86 R = 6.94 0.0002 0.00 Biological process Positive regulation of response GO:005171 C = 1717 O = 43 E = 2663 R = 1.82 0.0002 0.00 Biological process Positive regulation of response GO:00458 C = 1870 O = 43 E = 2663 R = 1.82 0.0002 0.00 Biological process Positive regulation of response GO:00458 C = 180 O = 17 E = 6.44 R = 2.84 0.0002 0.00 Biological process Immune effector process GO:000255 C = 120 O = 9 E = 2.12 R = 4.25 0.0002 0.00 Biological process Immune effector process GO:000255 C = 120 O = 9 E = 2.04 R = 4.42 0.0002 0.00 Biological process Immune effector process GO:000255 C = 120 O = 9 E = 0.15 R = 19.60 0.0004 0.00 Biological process Immune effector process GO:00025 C = 59 O = 6 E = 1.00 R = 0.00 0.0004 0.00 Biological process Immune response GO:00025 C = 59 O = 6 E = 1.00 R = 0.00 0.0004 0.00 or somatic recombination and presentation of account and the sentence of the 100 P = 0 = 100 R = 0.00 0.0005 0.00 Biological process Adaptive immune response based GO:00025 C = 59 O = 6 E = 1.05 R = 5.71 0.0006 0.00 antipue recombination of account process GO:000290 C = 3 O = 2 E = 0.05 R = 39.31 0.0008 0.00 antipue response Nature B C adaptive immune GO:00025 C = 62 O = 6 E = 1.05 R = 5.71 0.0006 0.00 Biological process Matter B cell appotein process GO:000290 C = 3 O = 2 E = 0.05 R = 39.31 0.0008 0.00 Biological process Matter B cell appotein process GO:000290 C = 3 O = 2 E = 0.05 R = 39.31 0.0008 0.00 Biological process Matter B cell appotein p	mediated by circulating		
Biological process Single-organism signaling GO:004470 C = 1510 O = 44 E = 25.61 R = 1.72 2.88 ⁻⁰⁵ O.0 Biological process Signal transduction GO:000716 C = 1308 O = 39 E = 22.18 R = 1.76 6.75 ⁻⁰³ 0.0 0.0 Biological process Positive regulation of GO:000377 C = 90 O = 6 E = 1.53 R = 5.24 0.0001 0.0 Biological process Cellular regionse GO:00274 C = 51 O = 6 E = 0.86 R = 6.94 0.0002 0.0 Biological process Lymphocyte mediated GO:000715 C = 1570 O = 43 E = 26.63 R = 1.82 0.0002 0.0 Biological process Cellular regionse GO:000715 C = 1570 O = 43 E = 2.64 R = 4.24 0.0002 0.0 Biological process Regulation of immune GO:000277 C = 120 O = 9 E = 2.12 R = 4.22 0.0002 0.0 Biological process Immune regionse GO:000277 C = 120 O = 9	immunoglobulin		
isgualing Biological process Signaling Biological process Signal increases GO:000716 C = 1308 C = 376 R = 1.76 R = 7.56 R = 2.51 0.00011 0.0 Biological process Positive regulation of GO:000277 C = 90 O = 8 E = 1.53 R = 5.24 0.00011 0.0 Biological process Califur response GO:000271 C = 1711 O = 46 E = 29.02 R = 1.59 0.0001 0.0 Biological process Cell communication GO:000271 C = 1711 O = 6 E = 0.86 R = 6.94 0.0002 0.0 Biological process Cell communication GO:000577 C = 1570 O = 43 E = 2.663 R = 1.82 0.0002 0.0 Biological process Regulation of immune GO:000577 C = 120 O = 9 E = 2.12 R = 4.42 0.0002 0.0 Biological process Immune response GO:000256 C = 120 O = 9 E = 2.04 R = 1.42 0.0002 0.0 Biological process	Biological process Signaling GO:002305 C = 1510 O = 44 E = 25.61 R = 1.72	2.68^{-05}	0.002
Biological process Signal transduction $GO:000716$ $C = 1308$ $O = 39$ $E = 2.218$ $R = 1.76$ $R = 2.51$ $O = 0.001$ $O = 0$ Biological process $GO:00037$ $C = 00$ $O = 8$ $E = 1.53$ $R = 5.24$ $O = 0.001$ $O = 0$ Biological proces $O = 0$ $O = 8$ $E = 1.53$ $R = 5.24$ $O = 0.001$ $O = 0$ Biological proces $O = 0$ $O = 8$ $E = 1.53$ $R = 5.24$ $O = 0.001$ $O = 0$ Biological proces $O = 0$ $O = 8$ $E = 0.86$ $R = 0.84$ $O = 0.002$ $O = 0$ Biological proces $O = 0$ $O = 8$ $E = 0.86$ $R = 0.84$ $O = 0.0002$ $O = 0$ Biological proces $O = 0$ $O = 0$ $O = 8$ $E = 0.86$ $R = 0.84$ $O = 0.0002$ $O = 0$ Biological proces $O = 0$	Biological process Single-organism GO:004470 C = 1510 O = 44 E = 25.61 R = 1.72	2.68^{-05}	0.002
Biological process Immune system process 00.000237 $C = 446$ $O = 19$ $E = 7.56$ $R = 2.51$ 0.0001 0.0 Biological process $Calkar response$ $GO.005077$ $C = 90$ $O = 8$ $E = 1.53$ $R = 5.24$ 0.0001 0.0 Immune response $GO.005171$ $C = 1711$ $O = 46$ $E = 29.02$ $R = 1.59$ $O.0001$ 0.0 Biological process $Calkar response$ $GO.000244$ $C = 51$ $O = 6$ $E = 0.86$ $R = 6.84$ 0.0002 0.0 Immunity $Calkar response$ $GO.000254$ $C = 51$ $O = 6$ $E = 0.86$ $R = 6.84$ 0.0002 0.0 Biological process $Calkar response$ $GO.000275$ $C = 1570$ $O = 43$ $E = 26.63$ $R = 1.62$ 0.0002 0.0 Biological process $Calkar response$ $GO.0002577$ $C = 125$ $O = 9$ $E = 2.12$ $R = 4.25$ 0.0002 0.0 Biological process $Regulation of response GO.000225 C = 120 O = 9 E = 2.04 R = 4.42 0.0002 0.0Biological process R response GO.000226 C = 10 O = 3 E = 0.17 R = 4.29 0.0002 0.0Biological process R adaptive immune response GO.000246 C = 59 O = 6 E = 1.00 R = 4.42 0.0002 0.0Biological process R adaptive immune response GO.000246 C = 59 O = 6 E = 1.00 R = 6.00 0.0004 0.00Biological process R calkar response GO.000246 C = 59 O = 6 E = 1.00 R = 6.00 0.0004 0.00Biological process R calkar response GO.000226 C = 59 O = 6 E = 1.00 R = 6.00 0.0005 0.0Biological process R calkar response GO.000226 C = 59 O = 6 E = 1.00 R = 7.69 O = 0.0005 $	signaling		
Biological process Positive regulation of moune response GO:005077 C = 90 O = 8 E = 1.53 R = 5.24 0.0001 0.00 Biological process Cellular response GO:005171 C = 1711 O = 46 E = 29.02 R = 1.59 0.0001 0.0 Biological process Lymphocyte modilated GO:005071 C = 1570 O = 6 E = 0.86 R = 6.94 0.0002 0.0 Biological process O estimulus GO:005077 C = 1570 O = 43 E = 26.63 R = 1.62 0.0002 0.0 Biological process Positive regulation of immune GO:005077 C = 125 O = 9 E = 2.12 R = 4.42 0.0002 0.0 Biological process Immune response GO:00025 C = 120 O = 9 E = 2.12 R = 4.42 0.0002 0.0 Biological process Immune response GO:00025 C = 120 O = 9 E = 0.15 R = 4.42 0.0002 0.0 Biological process B coll homeostals GO:000246 C = 59 O = 6 E = 0.17	Biological process Signal transduction GO:000716 C = 1308 O = 39 E = 22.18 R = 1.76	6.75^{-05}	0.0047
Immune response Biological process Cellular response GO:00171 C = 1711 O = 46 E = 29.02 R = 1.59 0.0001 0.0 Biological process Cell communication GO:000244 C = 51 O = 6 E = 0.86 R = 6.94 0.0002 0.0 Biological process Cell communication GO:000715 C = 1570 O = 43 E = 26.63 R = 1.62 0.0002 0.0 Biological process Regulation of mesponse GO:000275 C = 125 O = 9 E = 2.12 R = 4.42 0.0002 0.0 Biological process Immune regulation of response GO:000275 C = 120 O = 9 E = 2.12 R = 4.42 0.0002 0.0 Biological process Immune response GO:000246 C = 9 O = 3 E = 0.17 R = 4.25 0.0003 0.0 Biological process Adaptive immune response based GO:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GO:000247	Biological process Immune system process GO:000237 C = 446 O = 19 E = 7.56 R = 2.51	0.0001	0.0057
Biological process Caludar response to stimulus CO:005171 C = 1711 O = 46 E = 29.02 R = 1.59 0.001 0.0 Biological process Lymphocyte mediated immunity GO:000244 C = 51 O = 6 E = 0.86 R = 6.94 0.0002 0.0 Biological process Cell communication GO:000244 C = 1570 O = 43 E = 26.63 R = 1.62 0.0002 0.0 Biological process Positive regulation of immune response GO:0005077 C = 125 O = 9 E = 2.12 R = 4.25 0.0002 0.0 Biological process Immune effector process GO:000256 C = 120 O = 9 E = 2.12 R = 4.22 0.0002 0.0 Biological process Immune effector process GO:000266 C = 210 O = 3 E = 0.15 R = 14.66 0.0004 0.0 Biological process Adaptive immune response based GO:000246 C = 59 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Adaptive immune GO:000246 C = 30	Biological process Positive regulation of $GO:005077$ $C = 90$ $O = 8$ $E = 1.53$ $R = 5.24$	0.0001	0.0057
It is timulas Biological process Lymphocyte mediated G0:000244 C = 51 O = 6 E = 0.86 R = 6.94 0.0002 0.0 Biological process Cell communication G0:000715 C = 1570 O = 43 E = 6.44 R = 2.64 0.0002 0.0 Biological process Positive regulation of response G0:00027 C = 125 O = 9 E = 2.42 R = 4.25 0.0002 0.0 Biological process Immune regionse G0:000285 C = 120 O = 9 E = 2.04 R = 4.42 0.0002 0.0 Biological process Immune refeoro process G0:000285 C = 221 O = 12 E = 3.75 R = 3.20 0.0003 0.0 Biological process Manue response based G0:000246 C = 59 O = 6 E = 1.00 R = 6.00 0.0004 0.0 Biological process Antigen processing G0:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing G0:000225 C = 62	immune response		
Biological process Lymphocyte mediated immunity GO:000244 C = 51 O = 6 E = 0.86 R = 6.94 0.0002 0.0002 Biological process Cell communication GO:000715 C = 1570 O = 43 E = 26.63 R = 1.62 0.0002 0.0 Biological process Positive regulation of response GO:000777 C = 125 O = 9 E = 2.12 R = 4.25 0.0002 0.0 Biological process Immune effector process GO:000225 C = 120 O = 9 E = 2.12 R = 4.42 0.0002 0.0 Biological process B cell homeostasis GO:000265 C = 221 O = 12 E = 3.75 R = 4.42 0.0002 0.0 Biological process B cell homeostasis GO:000246 C = 90 O = 6 E = 1.00 R = 6.00 0.0004 0.0 Biological process Adaptive immune response based GO:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GO:000290 C = 342 O = 15	Biological process Cellular response GO:005171 C = 1711 O = 46 E = 29.02 R = 1.59	0.0001	0.0057
Immunity Biological process Cell communication G0:000715 C = 1570 O = 437 E = 26.63 R = 1.62 0.0002 0.0 Biological process Positive regulation of response G0:00077 C = 125 O = 9 E = 2.12 R = 4.25 0.0002 0.0 Biological process Immune fetetor process G0:000225 C = 120 O = 9 E = 2.04 R = 4.42 0.0002 0.0 Biological process Immune reflector process G0:000225 C = 120 O = 9 E = 2.04 R = 4.42 0.0003 0.0 Biological process Immune response G0:000245 C = 9 O = 3 E = 0.15 R = 19.66 0.0004 0.0 Biological process Adaptive immune response based G0:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing G0:000225 C = 62 O = 6 E = 1.05 R = 5.71 0.0008 0.0 Biological process Matrigen processing G0:000220 C =	to stimulus		
Biological process Cell communication GC:000715 C = 1570 O = 43 E = 6.63 R = 1.62 0.0002 0.00 Biological process Positive regulation of response GC:004858 C = 380 O = 77 E = 6.44 R = 2.64 0.0002 0.0 Biological process Regulation of immune GC:000225 C = 125 O = 9 E = 2.04 R = 4.42 0.0002 0.0 Biological process Immune effector process GC:000225 C = 120 O = 9 E = 2.04 R = 4.42 0.0002 0.0 Biological process Immune response GC:000246 C = 59 O = 3 E = 0.17 R = 19.66 0.0004 0.0 Biological process Antigen processing GC:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GC:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GC:000239 C = 342 O = 15 E = 5.80	Biological process Lymphocyte mediated $GO:000244$ $C = 51$ $O = 6$ $E = 0.86$ $R = 6.94$	0.0002	0.0086
Biological process Positive regulation of response to stimulus GO:004858 C = 380 O = 17 E = 6.44 R = 2.64 0.0002 0.002 Biological process Regulation of immune GO:005077 C = 125 O = 9 E = 2.12 R = 4.25 0.0002 0.00 Biological process Immune effector process GO:000695 C = 120 O = 9 E = 2.12 R = 4.42 0.0002 0.00 Biological process Immune effector process GO:00078 C = 9 O = 3 E = 0.15 R = 19.66 0.0004 0.0 Biological process A digen processing GO:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GO:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GO:00025 C = 62 O = 6 E = 1.05 R = 2.59 0.0005 0.0 Biological process Adaptive regulation of immune eceptors built from immunoglobulin superial minute GO:000290 C = 3 O = 2 E = 0.05 R = 39.31 0.0008	immunity		
bolicity in the second seco	Biological process Cell communication $GO:000715$ $C = 1570$ $O = 43$ $E = 26.63$ $R = 1.62$	0.0002	0.0086
Biological process Regulation of immune response GO:005077 C = 125 O = 9 E = 2.12 R = 4.25 0.0002 0.0017 Biological process Immune effector process GO:000225 C = 120 O = 9 E = 2.04 R = 4.42 0.0002 0.00 Biological process Biolinomeostasis GO:000246 C = 221 O = 12 E = 3.75 R = 3.20 0.0004 0.00 Biological process Adaptive immune response based GO:000246 C = 59 O = 6 E = 0.15 R = 1.66 0.0004 0.00 Biological process Adaptive immune response based GO:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.00 Biological process Antigen processing and presentation of exogeneous petitie antigen GO:000290 C = 342 O = 15 E = 5.80 R = 2.59 0.0005 0.00 Biological process Adaptive immune GO:000290 C = 3 O = 2 E = 0.05 R = 39.31 0.0008 0.00 Biological process Mature B cell apoptotic process GO:00	Biological process Positive regulation of response $GO:004858$ $C = 380$ $O = 17$ $E = 6.44$ $R = 2.64$	0.0002	0.0086
Tesponse Biological process Immune effector process GC:00025 C = 120 O = 9 E = 2.04 R = 4.42 0.0002 0.00 Biological process Immune response GC:000695 C = 221 O = 12 E = 3.75 R = 3.20 0.0003 0.00 Biological process B cell homeostasis GC:000246 C = 9 O = 3 E = 0.15 R = 19.66 0.0004 0.0 Biological process Adaptive immune response based GC:000246 C = 59 O = 6 E = 1.00 R = 6.00 0.0005 0.0 Biological process Antigen processing GC:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GC:000290 C = 342 O = 15 E = 5.80 R = 2.59 0.0005 0.0 Biological process Adaptive immune GO:000290 C = 3 O = 2 E = 0.05 R = 39.31 0.0008 0.0 Biological process Mature B cell apoptotic GO:000290 C =	to stimulus		
Biological process Immune effector process GC:000225 C = 120 $0 = 9$ E = 2.04 R = 4.42 0.0002 0.0003 0.00 Biological process Immune response GC:000695 C = 221 0 = 12 E = 3.75 R = 3.20 0.0004 0.00 Biological process Adaptive immune response based GC:000246 C = 9 0 = 3 E = 0.15 R = 19.66 0.0004 0.00 Biological process Adaptive immune response based GC:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.00 Biological process Antigen processing GC:000247 C = 10 O = 3 E = 0.17 R = 17.69 0.0005 0.0 Biological process Antigen processing GC:000225 C = 62 O = 6 E = 1.05 R = 5.71 0.0008 0.0 Biological process Mature B cell apoptotic process GC:000290 C = 3 O = 2 E = 0.05 R = 39.31 0.0008 0.0 Biological process Mature B cell apoptotic process GC:000290 C	Biological process Regulation of immune GO:005077 C = 125 O = 9 E = 2.12 R = 4.25	0.0002	0.0086
Biological processImmune responseGO:000695C = 221O = 12E = 3.75R = 3.200.00030.00Biological processB cell homeostasisGO:000178C = 9O = 3E = 0.15R = 19.660.00040.0Biological processAdaptive immune response basedGO:000246C = 59O = 6E = 1.00R = 6.000.00040.0Biological processAdaptive immune response basedGO:000247C = 10O = 3E = 0.17R = 17.690.00050.0Biological processAntigen processingGO:000247C = 10O = 3E = 0.17R = 17.690.00050.0Biological processAntigen processingGO:000295C = 62O = 6E = 1.05R = 2.590.00050.0Biological processResponse to lipidGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMatpitve immuneGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000295C = 66O = 6E = 1.12R = 5.360.00080.0Biological processRegulation of matureGO:000255C = 66O = 7E = 1.	response		
Biological processB cell homeostasisGO:000178C = 9O = 3E = 0.15R = 19.660.00040.0Biological processAdaptive immune response basedGO:000246C = 59O = 6E = 1.00R = 6.000.00040.0Biological processAdaptive immune globulin auperfamily domainsGO:000247C = 10O = 3E = 0.17R = 17.690.00050.0Biological processAntigen processing and presentation of exogenous peptide antigenGO:000247C = 10O = 3E = 0.17R = 17.690.00050.0Biological processResponse to lipidGO:000290C = 342O = 15E = 5.80R = 2.590.00050.0Biological processAdaptive immuneGO:000290C = 342O = 6E = 1.05R = 5.710.00080.0Biological processNegative regulation apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000295C = 66O = 6E = 1.12R = 5.360.00080.0Biological processPositive regulation of mmune system processGO:000255C = 66O = 7E = 1.56R = 4.49 </td <td>$Biological process \qquad Immune effector process \qquad GO:000225 \qquad C = 120 \qquad O = 9 \qquad E = 2.04 \qquad R = 4.42$</td> <td>0.0002</td> <td>0.0086</td>	$ Biological process \qquad Immune effector process \qquad GO:000225 \qquad C = 120 \qquad O = 9 \qquad E = 2.04 \qquad R = 4.42 $	0.0002	0.0086
Biological processAdaptive immune response based on somatic recombination of immune receptors built from immunoglobulin superfamily domainsGO:000246C = 59O = 6E = 1.00R = 6.000.00040.0Biological processAntigen processing antigen processingGO:000247C = 10O = 3E = 0.17R = 17.690.00050.0Biological processAntigen processing antigenGO:003399C = 342O = 15E = 5.80R = 2.590.00050.0Biological processResponse to lipidGO:003290C = 342O = 6E = 1.05R = 5.710.00060.0Biological processAdaptive immune responseGO:000225C = 62O = 6E = 1.05R = 5.710.00080.0Biological processNegative regulation of mature B cell apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of immune responseGO:000225C = 66O = 6E = 1.12R = 5.360.00080.0Biological processRegulation of immune responseGO:000225C = 66O = 6E = 1.12R = 5.360.00080.0Biological processRegulation of immune responseGO:000225C = 66O = 7E = 1.56R = 4.490.00090.0Biological processEpidermis developmentGO:0	Biological process Immune response GO:000695 C = 221 O = 12 E = 3.75 R = 3.20	0.0003	0.0124
biological process Activation of immune receptors built from immune graduation of and presentation of exogenous peptide antigen consists antigen consists and presentation of exogenous peptide antigen process and presentation of exogenous peptide antigen process and presentation of exogenous peptide and presentation of peptide and presentat	Biological process B cell homeostasis $GO:000178$ $C = 9$ $O = 3$ $E = 0.15$ $R = 19.66$	0.0004	0.0151
of immune receptors built from immunoglobulin superfamily domainsBiological processAntigen processing exogenous peptide antigenGO:000247 exogenous peptide antigenC = 10 P = 3 P = 2.59C = 17.69 P = 2.59R = 17.69 P = 0.00050.005 0.00Biological processResponse to lipidGO:000290 responseC = 342 P = 0.000225C = 62 P = 0.65C = 5.71 P = 0.00060.0006 P = 0.0006Biological processNegative regulation of mature B cell apoptotic processGO:000290 P = 0.000290C = 3 P = 2D = 2 P = 0.05 P = 0.05R = 39.31 P = 39.310.0008 P = 0.00080.00 P = 0.0008Biological processMature B cell apoptotic process P = 0.000290C = 3 P = 2D = 2 P = 0.05 P = 0.05R = 39.31 P = 39.310.0008 P = 0.00080.0008 P = 0.0008Biological processMature B cell apoptotic process P = 0.000290C = 3 P = 2D = 2 P = 0.05 P = 0.05R = 39.31 P = 39.310.0008 P = 0.00080.0008 P = 0.0008Biological processMature B cell apoptotic process P = 0.000290C = 3 P = 2D = 2 P = 0.05R = 5.36 P = 39.310.0008 P = 0.00080.0008 P = 0.0008Biological processActivation of immune responseGO:000290 P = 0.000290C = 3 P = 0.25P = 5.59 P = 3.47R = 4.49 P = 0.00080.0008 P = 0.0008Biological processEpidernis development responseGO:000268 P = 153C = 92 P = 0.97 P = 1.59R = 4.49 P = 0.001 P = 2.59	Biological process Adaptive immune response based $GO:000246$ $C = 59$ $O = 6$ $E = 1.00$ $R = 6.00$	0.0004	0.0151
from immunoglobulin superfamily domainsBiological processAntigen processing and presentation of exogenous peptide antigenGO:000247C = 10O = 3E = 0.17R = 17.690.00050.0Biological processResponse to lipidGO:003399C = 342O = 15E = 5.80R = 2.590.00050.0Biological processAdaptive immuneGO:000225C = 62O = 6E = 1.05R = 5.710.00080.0Biological processNegative regulationGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 5.360.00080.0Biological processActivation of immuneGO:000255C = 66O = 6E = 1.12R = 5.360.00080.0Biological processEpidermis developmentGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processProcessGO:004508C = 94O = 7E = 1.59R = 4.390.00	on somatic recombination		
Superfamily domainsBiological processAntigen processing and presentation of exogenous peptide antigenGO:000247 $C = 10$ $O = 3$ $E = 0.17$ $R = 17.69$ $O.0005$ $O.0005$ Biological processResponse to lipidGO:003399 $C = 342$ $O = 15$ $E = 5.80$ $R = 2.59$ $O.0005$ $O.0005$ Biological processAdaptive immuneGO:000225 $C = 62$ $O = 6$ $E = 1.05$ $R = 5.71$ $O.0006$ $O.0006$ Biological processNegative regulationGO:000290 $C = 3$ $O = 2$ $E = 0.05$ $R = 39.31$ $O.0008$ $O.0008$ Biological processMature B cell apoptotic process $O = 3$ $O = 2$ $E = 0.05$ $R = 39.31$ $O.0008$ $O.0008$ Biological processMature B cell apoptotic process $O = 3$ $O = 2$ $E = 0.05$ $R = 39.31$ $O.0008$ $O.0008$ Biological processRegulation of mature responseGO:000290 $C = 3$ $O = 2$ $E = 0.05$ $R = 39.31$ $O.0008$ $O.0008$ Biological processActivation of immune responseGO:000225 $C = 66$ $O = 6$ $E = 1.12$ $R = 5.36$ $O.0008$ $O.0009$ Biological processEpidermis developmentGO:000268 $C = 92$ $O = 7$ $E = 1.56$ $R = 4.49$ $O.0009$ $O.0009$ Biological processPositive regulation of responseGO:000268 $C = 153$ $O = 9$ $E = 2.59$ $R = 3.47$ $O.001$ $O.001$ Biological proc	of immune receptors built		
Biological process and presentation of exogenous peptide antigenAntigen processing exogenous peptide antigenGO:000247C = 10O = 3E = 0.17R = 17.690.00050.00Biological process Biological processResponse to lipidGO:003399C = 342O = 15E = 5.80R = 2.590.00050.00Biological processAdaptive immuneGO:000225C = 62O = 6E = 1.05R = 5.710.00060.00Biological processNegative regulation of mature B cell apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processMature B cell apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processRegulation of immuneGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processRegulation of immuneGO:000225C = 66O = 6E = 1.12R = 5.360.00090.0Biological processEpidermis developmentGO:000268C = 92O = 7E = 1.56R = 4.490.00090.0Biological processInmune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInmate immuneGO:004508C = 94O = 7E = 1.59R = 4.39<			
and presentation of exogenous peptide antigen Biological process Biological process Biol			
exogenous peptide antigenBiological processResponse to lipidGO:003399C = 342O = 15E = 5.80R = 2.590.00050.00Biological processAdaptive immuneGO:000225C = 62O = 6E = 1.05R = 5.710.00060.00Biological processNegative regulationGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processMature B cell apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processMature B cell apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processMature B cell apototic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processRegulation of immuneGO:000255C = 66O = 6E = 1.12R = 5.360.00080.00Biological processPositive regulation ofGO:000268C = 153O = 9E = 2.59R = 3.470.0010.00Immune system processImmune system processGO:004508C = 94O = 7E = 1.59R = 4.390.0010.00Biological process<		0.0005	0.0175
antigenBiological processResponse to lipidGO:003399C = 342O = 15E = 5.80R = 2.590.00050.0Biological processAdaptive immuneGO:000225C = 62O = 6E = 1.05R = 5.710.00060.0Biological processNegative regulationGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processof mature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processB cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processB cell apoptotic processC = 66O = 6E = 1.12R = 5.360.00080.0Biological processPointerinis developmentGO:000255C = 66O = 7E = 1.56R = 4.490.00090.0Biological processPointive regulation of responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processInnune system processGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Innune system processInnune responseFestoresFestoresFestoresFestoresFestoresFestores <td></td> <td></td> <td></td>			
Biological processResponse to lipidGO:003399C = 342O = 15E = 5.80R = 2.590.00050.0Biological processAdaptive immuneGO:000225C = 62O = 6E = 1.05R = 5.710.00060.0Biological processNegative regulationGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of immuneGO:000225C = 66O = 6E = 1.12R = 5.360.00080.0Biological processEpidermis developmentGO:000258C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0			
Biological processAdaptive immuneGO:000225C = 62O = 6E = 1.05R = 5.710.00060.0Biological processNegative regulationGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of immuneGO:000225C = 66O = 2E = 0.05R = 39.310.00080.0Biological processEpidermis developmentGO:000255C = 92O = 7E = 1.12R = 5.360.00090.0Biological processEpidermis developmentGO:000268C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation ofGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Immune system processInnate immuneGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0	•		
responseBiological processNegative regulation of mature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processB cell apoptotic processC = 66O = 6E = 1.12R = 5.360.00080.0Biological processEpidermis developmentGO:000255C = 66O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0			0.0175
Biological processNegative regulation of mature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processGO:000290C = 66O = 6E = 1.12R = 5.360.00080.0Biological processEpidermis developmentGO:000854C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:00268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0		0.0006	0.0202
of mature B cell apoptotic processBiological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processB cell apoptotic processGO:000225C = 66O = 6E = 1.12R = 5.360.00080.0Biological processActivation of immune responseGO:000854C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0			0 000 /
apoptotic processBiological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processB cell apoptotic processDecessC = 66O = 6E = 1.12R = 5.360.00080.00Biological processEpidermis developmentGO:000854C = 92O = 7E = 1.56R = 4.490.00090.00Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.00Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.00Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.00		0.0008	0.0234
Biological processMature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processRegulation of mature B cell apoptotic processGO:000290C = 3O = 2E = 0.05R = 39.310.00080.0Biological processB cell apoptotic processB cell apoptotic processGO:000225C = 66O = 6E = 1.12R = 5.360.00080.0Biological processEpidermis developmentGO:000854C = 92O = 7E = 1.56R = 4.490.00090.0Biological processEpidermis developmentGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0			
Biological processRegulation of matureGO:000290C = 3O = 2E = 0.05R = 39.310.00080.00Biological processB cell apoptotic processprocessGO:000225C = 66O = 6E = 1.12R = 5.360.00080.00Biological processActivation of immune responseGO:000254C = 92O = 7E = 1.56R = 4.490.00090.0Biological processEpidermis developmentGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0		0.0000	0.0004
B cell apoptotic processBiological processActivation of immune responseGO:000225C = 66O = 6E = 1.12R = 5.360.00080.0Biological processEpidermis developmentGO:000854C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0			0.0234
processBiological processActivation of immune responseGO:000225C = 66O = 6E = 1.12R = 5.360.00080.00Biological processEpidermis developmentGO:000854C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0		0.0008	0.0234
Biological processActivation of immune responseGO:000225C = 66O = 6E = 1.12R = 5.360.00080.0Biological processEpidermis developmentGO:000854C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0			
response Biological process Epidermis development GO:000854 C = 92 O = 7 E = 1.56 R = 4.49 0.0009 0.0 Biological process Positive regulation of immune system process GO:000268 C = 153 O = 9 E = 2.59 R = 3.47 0.001 0.0 Biological process Innate immune response GO:004508 C = 94 O = 7 E = 1.59 R = 4.39 0.001 0.0 Biological process Antigen processing and presentation of peptide GO:004800 C = 13 O = 3 E = 0.22 R = 13.61 0.0012 0.0		0.0000	0.0004
Biological processEpidermis developmentGO:000854C = 92O = 7E = 1.56R = 4.490.00090.0Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0		0.0008	0.0234
Biological processPositive regulation of immune system processGO:000268C = 153O = 9E = 2.59R = 3.470.0010.0Biological processInnate immune responseGO:004508C = 94O = 7E = 1.59R = 4.390.0010.0Biological processAntigen processing and presentation of peptideGO:004800C = 13O = 3E = 0.22R = 13.610.00120.0		0.0000	0.0055
immune system process Biological process Innate immune GO:004508 C = 94 O = 7 E = 1.59 R = 4.39 0.001 0.0 response Biological process Antigen processing and GO:004800 C = 13 O = 3 E = 0.22 R = 13.61 0.0012 0.0 presentation of peptide			0.0255
Biological process Innate immune GO:004508 C = 94 O = 7 E = 1.59 R = 4.39 0.001 0.00 Biological process Antigen processing and presentation of peptide GO:004800 C = 13 O = 3 E = 0.22 R = 13.61 0.0012 0.00		0.001	0.0267
response Biological process Antigen processing and GO:004800 C = 13 O = 3 E = 0.22 R = 13.61 0.0012 0.0 presentation of peptide 0.0012 <td></td> <td>0.001</td> <td>0.0007</td>		0.001	0.0007
$ \begin{array}{cccc} Biological \ process \\ presentation \ of \ peptide \end{array} \begin{array}{ccccc} GO:004800 & C = 13 & O = 3 & E = 0.22 & R = 13.61 & 0.0012 & 0.0 \\ \end{array} $		0.001	0.0267
presentation of peptide		0.0010	0.0004
		0.0012	0.0294
(Continued)			

GO terms	Term description	GO ID	Size of term	miRNA target	Expected	Enrichment ratio	Raw p value	FDR
	Antigen processing and	GO:001988	C = 13	O = 3	E = 0.22	R = 13.61	<i>p</i> value 0.0012	0.0294
Biological process	presentation of exogenous	GO.001966	0 - 13	0 = 3	E - 0.22	n - 13.01	0.0012	0.0294
	antigen							
Biological process	Leukocyte mediated	GO:000244	C = 71	O = 6	E = 1.20	R = 4.98	0.0012	0.0294
	immunity							
Biological process	Regulation of fibroblast	GO:004814	C = 29	O = 4	E = 0.49	R = 8.13	0.0013	0.0311
	proliferation							
Biological process	Multicellular organismal	GO:003250	C = 1895	O = 46	E = 32.14	R = 1.43	0.0017	0.0322
	process							
Biological process	Negative regulation	GO:000290	C = 4	O = 2	E = 0.07	R = 29.48	0.0017	0.0322
	of B cell apoptotic							
	process	0.0.00000	0 000	A 44	F (00	D 0.04	0.0004	0.000
Molecular function	Molecular transducer activity	GO:006008	C = 309	O = 14	E = 4.99	R = 2.81	0.0004	0.029
Molecular function	Signal transducer	GO:000487	C = 309	O = 14	E = 4.99	R = 2.81	0.0004	0.029
	activity	40.000401	0 000	0 14	L 4.00	11 2.01	0.0004	0.020
KEGG pathway	Staphylococcus aureus		C = 15	O = 6	E = 0.21	R = 28.48	2.97^{-08}	9.80^{-07}
	infection							
KEGG pathway	Complement and coagulation		C = 17	O = 6	E = 0.24	R = 25.13	7.19 ⁻⁰⁸	1.19^{-06}
	cascades							
KEGG pathway	Systemic lupus		C = 25	O = 5	E = 0.35	R = 14.24	2.10^{-05}	0.0002
	erythematosus							
KEGG pathway	p53 signaling pathway		C = 25	O = 3	E = 0.35	R = 8.55	0.0049	0.027
KEGG pathway	Antigen processing and		C = 25	O = 3	E = 0.35	R = 8.55	0.0049	0.027
	presentation							
KEGG pathway	Pathways in cancer		C = 142	O = 7	E = 1.99	R = 3.51	0.0037	0.027
KEGG pathway	MAPK signaling pathway		C = 123	O = 6	E = 1.73	R = 3.47	0.0075	0.0354
KEGG pathway	Pancreatic cancer		C = 34	O = 3	E = 0.48	R = 6.28	0.0117	0.0429
KEGG pathway	Colorectal cancer		C = 33	O = 3	E = 0.46	R = 6.47	0.0108	0.0429
KEGG pathway KEGG pathway	Lysosome		C = 70	O = 4	E = 0.98	R = 4.07	0.0165	0.0495
	Neuroactive ligand-receptor interaction		C = 72	O = 4	E = 1.01	R = 3.96	0.0182	0.0495
KEGG pathway	Chronic mveloid leukemia		C = 38	O = 3	E = 0.53	R = 5.62	0.0159	0.0495
KEGG pathway	Natural killer cell-mediated		C = 38 C = 41	0 = 3 0 = 3	E = 0.53 E = 0.58	R = 5.02 R = 5.21	0.0159	0.0495
	cytotoxicity		0 - 41	0 - 5	L - 0.30	n = 3.21	0.0135	0.0495

ciated with epileptogenesis (Vezzani et al., 2015). This is also supported by the identification of individual differentially expressed miRNAs (e.g., miR-146a-5p; Aronica et al., 2010) and mRNA targets (e.g., *CD74* and *C1r*, Teo and Wong, 2010; Zeis et al., 2016) involved in inflammation.

Worthy of note is the enrichment for genes in the MAPK signaling pathway. Enrichment within the MAPK cascade

has been reported during latency in the pilocarpine model in a hippocampal RNA expression study based on highthroughput RNA sequencing (Hansen et al., 2014). In particular, we found robust upregulation of two MAPKs, *Map3k14*, also called NIK, and *Map3k4*, also called MEKK4. *Map3k14*, a target of miR-7a-5p and miR-138-5p, mediates the neuron specific suppression of the nu-

			Size of	miRNA		Enrichment	Raw	
GO terms	Term description	GO ID	term	target	Expected	ratio	p value	FDR
Biological process	Negative regulation of DC differentiation	GO:2001198	C = 2	O = 2	E = 0	R = 576.79	2.88E-06	0.0005
Biological process	Regulation of DC differentiation	GO:2001199	C = 2	O = 2	E = 0	R = 576.79	2.88E-06	0.0005
Biological process	DC differentiation	GO:0097028	C = 13	O = 2	E = 0.02	R = 88.74	0.0002	0.0178
Biological process	Glial cell development	GO:0021782	C = 69	O = 3	E = 0.12	R = 25.08	0.0002	0.0178
Biological process	Regulation of glial cell proliferation	GO:0060251	C = 16	O = 2	E = 0.03	R = 72.10	0.0003	0.0214
Biological process	Glial cell proliferation	GO:0014009	C = 23	O = 2	E = 0.04	R = 50.16	0.0007	0.0356
Biological process	Regulation of immune system process	GO:0002682	C = 653	O = 6	E = 1.13	R = 5.30	0.0007	0.0356
Biological process	Response to wounding	GO:0009611	C = 692	O = 6	E = 1.20	R = 5	0.0009	0.0401
Biological process	Negative regulation of DNA binding	GO:0043392	C = 31	O = 2	E = 0.05	R = 37.21	0.0013	0.0498
Biological process	Oligodendrocyte development	GO:0014003	C = 32	O = 2	E = 0.06	R = 36.05	0.0014	0.0498
KEGG pathway	Hematopoietic cell lineage		C = 124	O = 3	E = 0.07	R = 40.29	5.70E-05	0.001
KEGG pathway	Lysosome		C = 79	O = 2	E = 0.05	R = 42.16	0.001	0.0015

Table 9. GO and KEGG enrichment results for the 29 predicted and anticorrelated mRNAs targets of five miRNAs differentially expressed in the chronic period

clear factor κ -B (*NF-kB*; Mao et al, 2016) that is upregulated in epilepsy patients (Teocchi et al., 2013) and in an experimental model of traumatic brain injury (Lipponen et al., 2016). *NF-kB* has been linked to traumatic brain injury relevant outcomes, including epileptogenesis and tissue repair, the hypothesis being that it plays an antiepileptogenic role (Lipponen et al., 2016). Thus, *Map3k14* activation may favor epileptogenesis and damage. The transcript of *Map3k4*, the other upregulated MAPK, is a target of four significantly downregulated miRNAs (namely, miR-92b-3p, miR-153-3p, miR-101a-3p, and miR-3573-3p). This enzyme activates the p38 and JNK pathways that are known to contribute to the apoptotic and inflammatory responses after kainate injection in mice (Yang et al., 1997; Jeon et al., 2000).

Activation of each of these suggested proepileptogenic kinase pathways might be counterbalanced by upregulation of inhibitors such as phosphatases. In our analysis, the protein tyrosine phosphatase, nonreceptor type 5 (Ptpn5), also called STEP, was found upregulated, and anticorrelated with the downregulated miR-150-5p and miR-383-5p. Contrary to Map3k4, Ptpn5 has been shown to inhibit p38 by selectively dephosphorylating its activation loop tyrosines and by sequestering it in the cytosol (Francis et al., 2014). Ptpn5 can also target the glutamate receptor subunits GluN2b and GluA2 leading to receptor internalization and decreased synaptic efficiency (Snyder et al., 2005; Xu et al., 2009). In addition, Ptpn5 inhibits the ERK2 pathway. Whereas p38 downstream molecules lead to the activation of neuroinflammation and apoptotic processes, the ERK2 cascade triggers neuronal differentiation and survival through the activation of the antiapoptotic gene *bcl-2* (Cruz and Cruz, 2007).

In addition to above, miRNAs such as miR-101a-3p, miR-551b-3p, and miR-139-5p may act together to modulate the expression of the predicted target syn2, a gene that is mutated in epileptic patients (Cavalleri et al., 2007). Synapsin 2 is a member of the synapsin family composed by synaptic vesicle phosphoproteins that modulate synaptic transmission and plasticity. Notably, Syn2-knockout mice show a decreased vesicle density at inhibitory synapses of DG GCs and are prone to epileptic seizures (Medrihan et al., 2013). Here, we found an inverse correlation of three downregulated miRNAs (miR-101a-3p, miR-551b-3p, and miR-139-5p) with the Syn2 mRNA, which levels are slightly increased suggesting that Syn2 phophoproteins, at this stage (i.e., epileptogenesis), are still able to control neuronal transmission at the DG synapses. Notably, Syn2 interacts with presynaptic Ca²⁺ channels to promote GABA asynchronous release (Medrihan et al., 2013), maintaining the tonic inhibition of excitatory neurons and contrasting the aberrant network synchronization that lead to seizures development in the chronic phase. These findings suggest an antiepileptogenic role of this inverse-correlation. The dentate cells, in this case, may slow down the miRNA levels to contrast the upcoming epileptogenic process.

High levels of miR-212-5p may favor the epileptogenic process through reduced expression of *Gabrd*. We found that the expression of the subunit δ of GABA_A receptors

was decreased in DG and inverse-correlated with the upregulated miR-212-5p. δ -Subunit-containing receptors are found in extrasynaptic and perisynaptic locations in hippocampal DG GCs (Wei et al., 2003). Because of their high affinity for GABA, they mediate tonic GABA_A inhibition (Stell et al., 2003). Therefore, a decrease in GABA_A receptor δ -subunits may impair tonic GABA inhibition, contributing to GCs hyperexcitation and seizures onset.

Chronic epilepsy

Our exploration of the chronic stage of epilepsy was more limited than that for epileptogenesis as anticorrelations (inferred via statistically significant gene and miRNA fold changes in response to the disease) with miRNA targets could be evaluated only for the amygdala stimulation model. This analysis highlighted the downregulation of five miRNAs and the upregulation of several mRNAs targets, and identified genes enriched in GO terms related to glial cells and dendritic cells (DCs). Whereas the proliferation of glia cells and their contribution to neuroinflammation and hyperexcitability in chronic epilepsy are well recognized (Devinsky et al., 2013; Robel and Sontheimer, 2016), the role of DCs in the context of epilepsy remains elusive. It can be hypothesized that DCs might be involved in epilepsy by maintaining a chronic inflammatory response (Ludewig et al., 2016).

Among the list of mRNAs predicted targets anticorrelated with significantly differentially expressed miRNAs in the chronic stage of epilepsy, of note is the glutamate ionotropic receptor δ type 2 (*Grid2*), which is anticorrelated with miR-130a-3p, miR-148b-3p, and miR-551b-3p. The involvement of ionotropic glutamate receptors in epileptic hyperexcitability is well established, but not much is known specifically on glutamate ionotropic type δ receptors or the regulation of this process. Further studies are needed to establish a role of these receptors in hippocampus and more specifically in epilepsy.

Limitations

The purpose (and, in our view, the strength) of this work was to maximize information from underpowered individual studies, increasing power and allowing the identification of a set of miRNAs that, being significantly and similarly dysregulated in multiple experimental models, may be related to the disease rather than specific to a particular model. The study, however, also has limitations that should be taken into account.

A technical limitation is that the comparison of datasets in which tissue was obtained through different methods and that used different microarray platforms may have led to some miRNAs being detected in one experimental model and not in another, due to technical differences related to the assay system. Therefore, we cannot exclude the possibility that additional miRNAs were significantly dys-regulated.

Other limitations refer to biological aspects. First, miRNAs are only one mechanism of regulation of gene expression. Other changes may occur depending on other epigenetic mechanisms (histone modifications, DNA methylations) or changes in transcription factors. Second, this analysis has been conducted on one specific hippocampal

New Research 21 of 22

eNeuro

subarea, the DG, enriched in a specific cell population, the GCs. Other brain areas and cell populations may be equally or even more important in epileptogenesis. Third, due to limitation in the availability of datasets, we analyzed data from a single time point in all models, but epileptogenesis may develop differently in different models. Finally, given the lack of miRNA and mRNA datasets in epileptic patients matched with valid controls, we could not verify whether the miRNA-mRNA interactions identified in rats may be relevant for the human disease.

In addition, the comparison with mRNA datasets should be viewed as a secondary outcome of the study and considered with caution. In this study, this comparison is primarily based on the assumption that miRNAs decrease levels of their mRNA targets (Guo et al., 2010) and, therefore, that target mRNAs will undergo changes in anticorrelation with those of miRNAs. Although this is the best characterized mechanism of miRNA action, it is becoming evident that miRNAs also have specific nuclear functions, including transcriptional control of gene expression and regulation of alternative splicing (Catalanotto et al., 2016), which may not lead to anticorrelation between miRNA and mRNA levels. Therefore, we also performed a further analysis of direct correlation between miRNAs and mRNAs. These analyses now require verification and further studies to establish the exact patterns of interaction between miRNAs and mRNAs in the epileptic tissue and their functional impact on epileptogenesis and maintenance of an epileptic condition.

Conclusions

The present meta-analysis identified many significantly differentially expressed miRNAs in epileptogenesis and chronic epilepsy, several of which were not uncovered in the individual studies, highlighting the additional information that can be gained by meta-analysis. Our results also highlight the added value of meta-analysis of existing data and so avoid unnecessary animal experimentation to generate new hypotheses on miRNAs involved in epileptogenesis and chronic epilepsy. Our results highlight a possible key role for a few miRNAs that are worthy of further investigation. As it may be expected, however, the number and heterogeneity of mRNAs identified by this meta-analysis suggest that therapies focused on a single miRNA target may be not sufficient to reverse or ameliorate the epileptogenic process.

References

- Aronica E, Fluiter K, Iyer A, Zurolo E, Vreijling J, van Vliet EA, Baayen JC, Gorter JA (2010) Expression pattern of miR-146a, an inflammation-associated microRNA, in experimental and human temporal lobe epilepsy. Eur J Neurosci 31:1100–1107. CrossRef
- Bartel DP (2004) MicroRNAs: genomics, biogenesis, mechanism, and function. Cell 116:281–297. Medline
- Berben L, Sereika SM, Engberg S (2012) Effect size estimation: methods and examples. Inter J Nurs Stud 49:1039–1047. Cross-Ref Medline
- Bot AM, Dębski KJ, Lukasiuk K (2013) Alterations in miRNA levels in the dentate gyrus in epileptic rats. PLoS One 8:e76051. CrossRef Medline
- Brennan GP, Dey D, Chen Y, Patterson KP, Magnetta EJ, Hall AM, Dube CM, Mei YT, Baram TZ (2016) Dual and opposing roles of microRNA-124 in epilepsy are mediated through inflammatory and NRSF-dependent gene networks. Cell Rep 14:2402–2412. Cross-Ref Medline

- Catalanotto C, Cogoni C, Zardo G (2016) MicroRNA in control of gene expression: an overview of nuclear functions. Int J Mol Sci 17:pii:E1712. CrossRef
- Cattani AA, Allene C, Seifert V, Rosenow F, Henshall DC, Freiman TM (2016) Involvement of microRNAs in epileptogenesis. Epilepsia 57:1015–1026. CrossRef Medline
- Cavalleri GL, Weale ME, Shianna KV, Singh R, Lynch JM, Grinton B, Szoeke C, Murphy K, Kinirons P, O'Rourke D, Ge D, Depondt C, Claeys KG, Pandolfo M, Gumbs C, Walley N, McNamara J, Mulley JC, Linney KN, Sheffield LJ, et al. (2007) Multicentre search for genetic susceptibility loci in sporadic epilepsy syndrome and seizure types: a case-control study. Lancet Neurol 6:970–980. CrossRef Medline
- Chevaleyre V, Siegelbaum SA (2010) Strong CA2 pyramidal neuron synapses define a powerful disynaptic cortico-hippocampal loop. Neuron 66:560–572. CrossRef Medline
- Cruz CD, Cruz F (2007) The ERK 1 and 2 pathway in the nervous system: from basic aspects to possible clinical applications in pain and visceral dysfunction. Curr Neuropharmacol 5:244–252. Cross-Ref Medline
- DerSimonian R, Laird N (1986) Meta-analysis in clinical trials. Control Clini Trials 7:177–188. Medline
- DerSimonian R, Kacker R (2007) Random-effects model for metaanalysis of clinical trials: an update. Contemp Clin Trials 28:105– 114. CrossRef Medline
- Devinsky O, Vezzani A, Najjar S, De Lanerolle NC, Rogawski MA (2013) Glia and epilepsy: excitability and inflammation. Trends Neurosci 36:174–184. CrossRef Medline
- Dingledine R, Coulter DA, Fritsch B, Gorter JA, Lelutiu N, McNamara J, Nadler JV, Pitkänen A, Rogawski MA, Skene P, Sloviter RS, Wang Y, Wadman WJ, Wasterlain C, Roopra A (2017) Transcriptional profile of dentate granule cells in four rat epilepsy models. Sci Data 4:170061. CrossRef Medline
- Dudek FE, Sutula TP (2007) Epileptogenesis in the dentate gyrus: a critical perspective. Prog Brain Res 163:755–773. CrossRef Medline
- Dweep H, Sticht C, Pandey P, Gretz N (2011) miRWalk–database: prediction of possible miRNA binding sites by "walking" the genes of three genomes. J Biomed Inform 44:839–847. CrossRef
- Dweep H, Gretz N, Sticht C (2014) miRWalk database for miRNA-target interactions. Methods Mol Biol 1182:289–305. CrossRef Medline
- Ebert MS, Sharp PA (2012) Roles for microRNAs in conferring robustness to biological processes. Cell 149:515–524. CrossRef Medline
- Francis DM, Koveal D, Tortajada A, Page R, Peti W (2014) Interaction of kinase-interaction-motif protein tyrosine phosphatases with the mitogen-activated protein kinase ERK2. PLoS One 9:e91934. CrossRef Medline
- Gorter JA, Iyer A, White I, Colzi A, van Vliet EA, Sisodiya S, Aronica E (2014) Hippocampal subregion-specific microRNA expression during epileptogenesis in experimental temporal lobe epilepsy. Neurobiol Dis 62:508–520. CrossRef Medline
- Grimson A, Farh KK, Johnston WK, Garrett-Engele P, Lim LP, Bartel DP (2007) MicroRNA targeting specificity in mammals: determinants beyond seed pairing. Mol Cell 27:91–105. CrossRef Medline
- Guo H, Ingolia NT, Weissman JS, Bartel DP (2010) Mammalian microRNAs predominantly act to decrease target mRNA levels. Nature 466:835–840. CrossRef Medline
- Hansen KF, Sakamoto K, Pelz C, Impey S, Obrietan K (2014) Profiling status epilepticus-induced changes in hippocampal RNA expression using high-throughput RNA sequencing. Sci Rep 4:6930. CrossRef Medline
- Hauser WA, Annegers JF, Kurland LT (1993) Incidence of epilepsy and unprovoked seizures in Rochester, Minnesota: 1935-1984. Epilepsia 34:453–468. Medline
- Henshall DC, Hamer HM, Pasterkamp RJ, Goldstein DB, Kjems J, Prehn JH, Schorge S, Lamottke K, Rosenow F (2016) MicroRNAs in epilepsy: pathophysiology and clinical utility. Lancet Neurol 15:1368–1376. CrossRef Medline
- Higgins JP, Thompson SG, Deeks JJ, Altman DG (2003) Measuring inconsistency in meta-analyses. BMJ 327:557–560. CrossRef Medline

- Ioannidis JP, Patsopoulos NA, Evangelou E (2007) Heterogeneity in meta-analyses of genome-wide association investigations. PLoS One 2:e841. CrossRef Medline
- Jeon SH, Kim YS, Bae CD, Park JB (2000) Activation of JNK and p38 in rat hippocampus after kainic acid induced seizure. Exp Mol Med 32:227–230. CrossRef Medline
- Jimenez-Mateos EM, Engel T, Merino-Serrais P, McKiernan RC, Tanaka K, Mouri G, Sano T, O'Tuathaigh C, Waddington JL, Prenter S, Delanty N, Farrell MA, O'Brien DF, Conroy RM, Stallings RL, DeFelipe J, Henshall DC (2012) Silencing microRNA-134 produces neuroprotective and prolonged seizure-suppressive effects. Nat Med 18:1087–1094. CrossRef Medline
- Jinek M, Doudna JA (2009) A three-dimensional view of the molecular machinery of RNA interference. Nature 457:405–412. Cross-Ref Medline
- Karnati HK, Panigrahi MK, Gutti RK, Greig NH, Tamargo IA (2015) miRNAs: key players in neurodegenerative disorders and epilepsy. J Alzheimers Dis 48:563–580. CrossRef Medline
- Krook-Magnuson E, Armstrong C, Bui A, Lew S, Oijala M, Soltesz I (2015) *In vivo* evaluation of the dentate gate theory in epilepsy. J Physiol 593:2379–2388. CrossRef Medline
- Lipponen A, Paananen J, Puhakka N, Pitkänen A (2016) Analysis of post-traumatic brain injury gene expression signature reveals tubulins, Nfe2l2, Nfkb, Cd44, and S100a4 as treatment targets. Sci Rep 6:31570. CrossRef Medline
- Ludewig P, Gallizioli M, Urra X, Behr S, Brait VH, Gelderblom M, Magnus T, Planas AM (2016) Dendritic cells in brain diseases. Biochim Biophys Acta 1862:352–367. CrossRef Medline
- Lukasiuk K, Pitkänen A (2004) Large-scale analysis of gene expression in epilepsy research: is synthesis already possible? Neurochem Res 29:1169–1178. Medline
- Mao X, Phanavanh B, Hamdan H, Moerman-Herzog AM, Barger SW (2016) NFκB-inducing kinase inhibits NFκB activity specifically in neurons of the CNS. J Neurochem 137:154–163. CrossRef
- Medrihan L, Cesca F, Raimondi A, Lignani G, Baldelli P, Benfenati F (2013) Synapsin II desynchronizes neurotransmitter release at inhibitory synapses by interacting with presynaptic calcium channels. Nat Commun 4:1512. CrossRef Medline
- Peng Z, Houser CR (2005) Temporal patterns of fos expression in the dentate gyrus after spontaneous seizures in a mouse model of temporal lobe epilepsy. J Neurosci 25:7210–7220. CrossRef
- Pitkänen A, Lukasiuk K (2011) Mechanisms of epileptogenesis and potential treatment targets. The Lancet Neurol 10:173–186. Cross-Ref Medline
- Polit DF, Beck CT (2004) Nursing research. Principles and methods. Philadelphia: Lippincott Williams & Wilkins.
- Rajman M, Metge F, Fiore R, Khudayberdiev S, Aksoy-Aksel A, Bicker S, Ruedell Reschke C, Raoof R, Brennan GP, Delanty N, Farrell MA, O'Brien DF, Bauer S, Norwood B, Veno MT, Krüger M, Braun T, Kjems J, Rosenow F, Henshall DC, et al. (2017) A microRNA-129-5p/Rbfox crosstalk coordinates homeostatic downscaling of excitatory synapses. EMBO J 36:1770–1787. CrossRef Medline
- Ramasamy A, Mondry A, Holmes CC, Altman DG (2008) Key issues in conducting a meta-analysis of gene expression microarray datasets. PLoS Med 5:e184. CrossRef Medline
- Robel S, Sontheimer H (2016) Glia as drivers of abnormal neuronal activity. Nat Neurosci 19:28–33. CrossRef Medline
- Roncon P, Soukupovà M, Binaschi A, Falcicchia C, Zucchini S, Ferracin M, Langley SR, Petretto E, Johnson MR, Marucci G, Michelucci R, Rubboli G, Simonato M (2015) MicroRNA profiles in hippocampal granule cells and plasma of rats with pilocarpineinduced epilepsy-comparison with human epileptic samples. Sci Rep 5:14143. CrossRef Medline
- Roncon P, Zucchini S, Ferracin M, Marucci G, Giulioni M, Michelucci R, Rubboli G, Simonato M (2016) Is autopsy tissue a valid control for epilepsy surgery tissue in microRNA studies? Epilepsia Open 2:90–95. CrossRef
- Shin C, Nam JW, Farh KK, Chiang HR, Shkumatava A, Bartel DP (2010) Expanding the microRNA targeting code: functional sites with centered pairing. Mol Cell 38:789–802. CrossRef Medline

- Shkumatava A, Stark A, Sive H, Bartel DP (2009) Coherent but overlapping expression of microRNAs and their targets during vertebrate development. Genes Dev 23:466–481. CrossRef Medline
- Simonato M, Löscher W, Cole AJ, Dudek FE, Engel J Jr, Kaminski RM, Loeb JA, Scharfman H, Staley KJ, Velíšek L, Klitgaard H (2012) Finding a better drug for epilepsy: preclinical screening strategies and experimental trial design. Epilepsia 53:1860–1867. CrossRef
- Simonato M, French JA, Galanopoulou AS, O'Brien TJ (2013) Issues for new antiepilepsy drug development. Curr Opin Neurol 26:195– 200. CrossRef Medline
- Simonato M, Brooks-Kayal AR, Engel J Jr, Galanopoulou AS, Jensen FE, Moshé SL, O'Brien TJ, Pitkanen A, Wilcox KS, French JA (2014) The challenge and promise of anti-epileptic therapy development in animal models. Lancet Neurol 13:949–960. CrossRef
- Snyder EM, Nong Y, Almeida CG, Paul S, Moran T, Choi EY, Nairn AC, Salter MW, Lombroso PJ, Gouras GK, Greengard P (2005) Regulation of NMDA receptor trafficking by amyloid-beta. Nat Neurosci 8:1051–1058. CrossRef Medline
- Stell BM, Brickley SG, Tang CY, Farrant M, Mody I (2003) Neuroactive steroids reduce neuronal excitability by selectively enhancing tonic inhibition mediated by delta subunit-containing GABAA receptors. Proc Natl Acad Sci USA 100:14439–14444. CrossRef Medline
- Tan CL, Plotkin JL, Venø MT, von Schimmelmann M, Feinberg P, Mann S, Handler A, Kjems J, Surmeier DJ, O'Carroll D, Greengard P, Schaefer A (2013) MicroRNA-128 governs neuronal excitability and motor behavior in mice. Science 342:1254–1258. CrossRef Medline
- Teo BH, Wong SH (2010) MHC class II-associated invariant chain (li) modulates dendritic cells-derived microvesicles (DCMV)-mediated activation of microglia. Biochem Biophys Res Commun 400:673– 678. CrossRef
- Teocchi MA, Ferreira AÉ, da Luz de Oliveira EP, Tedeschi H, D'Souza-Li L (2013) Hippocampal gene expression dysregulation of Klotho, nuclear factor kappa B and tumor necrosis factor in temporal lobe epilepsy patients. J Neuroinflammation 10:53. CrossRef Medline
- Vezzani A, Lang B, Aronica E (2015) Immunity and inflammation in epilepsy. Cold Spring Harb Perspect Med 6:a022699. CrossRef Medline
- Wang J, Duncan D, Shi Z, Zhang B (2013) WEB-based GEne SeT AnaLysis Toolkit (WebGestalt): update. Nucleic Acids Res 41: W77–W83. CrossRef Medline
- Wei W, Zhang N, Peng Z, Houser CR, Mody I (2003) Perisynaptic localization of delta subunit-containing GABA(A) receptors and their activation by GABA spillover in the mouse dentate gyrus. J Neurosci 23:10650–10661. Medline
- Xi ZQ, Xiao F, Yuan J, Wang XF, Wang L, Quan FY, Liu GW (2009) Gene expression analysis on anterior temporal neocortex of patients with intractable epilepsy. Synapse 63:1017–1028. CrossRef Medline
- Xu J, Kurup P, Zhang Y, Goebel-Goody SM, Wu PH, Hawasli AH, Baum ML, Bibb JA, Lombroso PJ (2009) Extrasynaptic NMDA receptors couple preferentially to excitotoxicity via calpain-mediated cleavage of STEP. J Neurosci 29:9330–9343. CrossRef Medline
- Yang DD, Kuan CY, Whitmarsh AJ, Rincón M, Zheng TS, Davis RJ, Rakic P, Flavell RA (1997) Absence of excitotoxicity-induced apoptosis in the hippocampus of mice lacking the Jnk3 gene. Nature 389:865–870. CrossRef Medline
- Yang Z, Chen Y, Fu Y, Yang Y, Zhang Y, Chen Y, Li D (2014) Meta-analysis of differentially expressed genes in osteosarcoma based on gene expression data. BMC Med Genet 15:80. CrossRef Medline
- Yuzaki M (2017) The C1q complement family of synaptic organizers: not just complementary. Curr Opin Neurobiol 45:9–15. CrossRef Medline
- Zeis T, Enz L, Schaeren-Wiemers N (2016) The immunomodulatory oligodendrocyte. Brain Res 1641:139–148. CrossRef
- Zhang B, Kirov S, Snoddy J (2005) WebGestalt: an integrated system for exploring gene sets in various biological contexts. Nucleic Acids Res 33:W741–W748. CrossRef Medline