Dysregulated miRNA biogenesis downstream of cellular stress and ALS-causing mutations: a new mechanism for ALS

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Abstract

Interest in RNA dysfunction in amyotrophic lateral sclerosis (ALS) recently aroused upon discovering causative mutations in RNA-binding protein genes. Here, we show that extensive down-regulation of miRNA levels is a common molecular denominator for multiple forms of human ALS. We further demonstrate that pathogenic ALS-causing mutations are sufficient to inhibit miRNA biogenesis at the Dicing step. Abnormalities of the stress response are involved in the pathogenesis of neurodegeneration, including ALS. Accordingly, we describe a novel mechanism for modulating microRNA biogenesis under stress, involving stress granule formation and re-organization of DICER and AGO2 protein interactions with their partners. In line with this observation, enhancing DICER activity by a small molecule, enoxacin, is beneficial for neuromuscular function in two independent ALS mouse models. Characterizing microRNA biogenesis downstream of the stress response ties seemingly disparate pathways in neurodegeneration and further suggests that DICER and miRNAs affect neuronal integrity and are possible therapeutic targets.

Keywords microRNA, ALS, stress, neurodegeneration, DICER

Subject Categories Neuroscience; RNA Biology

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Introduction

Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease of the human motor neuron system. Mutations in genes encoding RNA-binding proteins were recently identified as causative in human ALS. These include TAR DNA-binding protein 43 (TDP-43) (Kabashi et al, 2008; Sreedharan et al, 2008; Liu-Yesucevitz et al, 2010; McDonald et al, 2011), fused in sarcoma (FUS) (Kwiatkowski et al, 2009; Vance et al, 2009; Bosco et al, 2010; Gal et al, 2010) and heterogeneous nuclear ribonucleoprotein A1 (hnRNP A1) (Kim et al, 2013).

Stress granules (SGs) are cytoplasmic sites for modulating mRNA translation and form in response to cellular stress (Anderson & Kedersha, 2008, 2009; Buchan & Parker, 2009). Intriguingly, among the RNA-binding proteins that are recruited into SGs, several are mutated in ALS or the related disorder fronto-temporal dementia (FTD), for example, TDP-43 and FUS.

Stress and SGs are thought to be involved in the pathogenesis of ALS, and SGs are observed in pathologic ALS/FTD specimen (Neumann et al, 2007; Volkenling et al, 2009; Bosco et al, 2010; Dormann et al, 2010; Gal et al, 2010; Liu-Yesucevitz et al, 2010; Dewey et al, 2011; McDonald et al, 2011; Meyerowitz et al, 2011; Aulas et al, 2012; Baron et al, 2013; Daigle et al, 2013; Kim et al, 2013; Vance et al, 2013; reviewed in Dewey et al, 2012; Emde & Hornstein, 2014; Li et al, 2013b; Ling et al, 2013). SG formation is initiated by the phosphorylation of eukaryotic translation initiation factor 2 alpha (eIF2A) on serine-51 or by the expression of several aggregation-prone proteins that are capable of initiating SG
nucleation (Kedersha et al., 1999). Accordingly, modulation of EIF2A phosphorylation mitigates neurotoxicity in animal models of ALS (Saxena et al., 2009; Kim et al., 2014).

miRNAs (miRNAs) are small endogenous RNAs. Initial processing of primary miRNAs by Drosha/Dgcr8 (Gregory et al., 2004) yields an intermediate pre-miRNA that is further processed by the 500-kDa DICER multi-protein complex to form a mature miRNA. This complex contains DICER 1, ribonuclease type III (DICER) and its partners, Argonaute RISC catalytic component 2 (AGO2), TAR (HIV-1) RNA-binding protein 2 (also known as TRBP) and protein kinase, interferon-inducible double-stranded RNA-dependent activator (PACT) (Chendrimada et al., 2005; Haase et al., 2005; Lee et al., 2006). DICER activity is controlled by its core partners (Ma et al., 2008; Shan et al., 2008; Li et al., 2012; Lee et al., 2013) and the mature miRNA is loaded onto AGO2, creating a functional RNA-induced silencing complex (RISC).

DICER and its co-factors participate at multiple levels of the stress response (Emde & Hornstein, 2014). Thus, DICER contributes to stress resistance in selected tissues and its deficiency reduces stress tolerance (Mori et al., 2012). In addition, various stress-signaling cascades control AGO2 post-translational modifications, its translocation into SGs and its activity (Leung et al., 2006; Qi et al., 2008; Zeng et al., 2008; Shen et al., 2013).

We previously demonstrated that loss of miRNA biogenesis is sufficient to cause spinal motor neuron degeneration in vivo (Haramati et al., 2010). In the current work, we demonstrate that reduction in miRNA levels is a common molecular denominator for multiple forms of familial and sporadic human ALS and that enhancement of DICER activity is beneficial in vivo in two independent ALS mouse models. The observations establish a novel pathway downstream of the stress response that controls DICER complex activity. The regulation of DICER involves stress granule assembly and dynamic interactions with SG proteins. This new mechanism for the control of miRNA biogenesis offers novel therapeutic targets for ALS.

**Results**

**miRNAs are down-regulated in motor neurons of human ALS patients**

We recently demonstrated that conditional inactivation of DICER in motor neurons is sufficient to cause spinal motor neuron degeneration in vivo (Haramati et al., 2010). Additional works from other groups further suggest the involvement of miRNAs in ALS pathogenesis (Williams et al., 2009; Buratti et al., 2010; Kawahara & Mieda-Sato, 2012; Morlando et al., 2012; Droppelmann et al., 2013; Freischmidt et al., 2013; Koval et al., 2013; Zhang et al., 2013). To pursue this, we examined miRNA expression in lumbar motor neurons of sporadic ALS (sALS) spinal cords isolated by laser capture microdissection (LCM). Global down-regulation of miRNAs was observed in motor neurons. The observation seemed specific to motor neurons since RNA extracted from surrounding, motor neuron-depleted ventral-horn tissue of sALS patients or from neurons of Clarke’s column (non-motor neuron) isolated by LCM within the same nervous systems did not exhibit such global changes in miRNA expression (Fig 1A–C). The observations of miRNA down-regulation were substantiated by miRNA in situ hybridization, which revealed down-regulation of miR-9 and miR-124 in patient tissue, relative to control tissue and to the hybridization signal of U6 RNA (Fig 1D). We next investigated whether miRNAs display a similar profile in fALS cases. Indeed, analyzing two nervous systems carrying familial SOD1 A4V mutation, revealed similar down-regulation of miRNAs (Fig 1E and F) that correlated with miRNA changes, observed in sALS (Pearson’s correlation coefficient = 0.3). Our data suggest that changes in miRNA expression, which were previously reported in ALS spinal cord extracts (Campos-Melo et al., 2013), are primarily due to changes in miRNA expression in motor neurons. Attempts to measure pre-miRNAs in human LCM samples were unsuccessful, probably due to the low abundance of these intermediate precursors. Noteworthy, expression and splicing data for changes in miRNAs encoding miRNA biogenesis factors from the relevant ALS patients did not reveal any change, relative to controls (Rabin et al., 2009). Additionally, 27 long ncRNAs tested were similarly expressed in another cohort of 12 non-ALS and 12 sporadic ALS nervous systems, with Pearson’s correlation of 0.9882. Since miRNAs are globally down-regulated in sALS cases that are genetically unrelated, reduced miRNA expression in motor neurons appears to be a common denominator of various forms of ALS.

**Cellular stress affects pre-miRNA processing**

Stress pathways are thought to control miRNA activity (Leung & Sharp, 2010; Mendell & Olson, 2012; Emde & Hornstein, 2014), but the idea that cellular stress might broadly regulate miRNA biogenesis has not been tested. We sought to do so by the measurement of the relative levels of mature miRNAs and pre-miRNAs under several stress conditions. We further defined an “inhibition score” that approximates a value of one, when DICER processes pre-miRNAs effectively, as in wild-type conditions, and increases >1, when miRNA biogenesis is inhibited, relative to control (Fig 2A).

We chose representative miRNAs that are expressed in human and mouse motor neurons including miR-10b, miR-30a and miR-103 (abundantly expressed in isolated mouse motor neurons); miR-218, miR-30c, miR-138 and let-7b (highly expressed in human motor neurons); miR-132, let-7c and let-7d (neuronal, mid-range in human); and miR-143 and let-7a (low expression in human neurons).

We triggered cellular stress in the hybrid motor neuron cell line NSC-34; using thapsigargin induced markers of ER stress (Appendix Fig S1A), down-regulated mature miRNA levels and up-regulated pre-miRNA levels, relative to control, demonstrating the inhibition of miRNA biogenesis (Fig 2B–D). In addition, application of oxidative stressors, paraquat or sodium arsenite, led to the inhibition of miRNA biogenesis (Fig 2E–J, Appendix Fig S1B).

To test whether changes in DICER or its co-factors are down-regulated as a result of stress, we measured protein levels by Western blot analysis. The expression levels of the four DICER complex proteins were not down-regulated by two independent stressors (thapsigargin, paraquat). DICER and AGO2 levels were not reduced by sodium arsenite as well, but PACT and TRBP levels were reduced by 1/3 (Appendix Fig S2 related to Fig 2). We conclude that alteration of DICER activity may occur even without reduction in DICER complex components.
Expression of ALS-causing genes inhibits pre-miRNA processing

To assess whether ALS-causing genes affect miRNA biogenesis similar to chemical stressors, we transfected NSC-34 cells with vectors of wild-type FUS, TDP-43 and SOD1 or mutant forms of the same genes, specifically FUS R495X (Bosco et al., 2010; Waibel et al., 2010), TDP-43 A315T (Gitcho et al., 2008) and SOD1 G93A (Rosen et al., 1993) (Fig 3A). We observed inhibition in biogenesis of many of the miRNA species tested, relative to the corresponding control sample values, after overexpression of wild-type or mutant FUS and TDP-43 (Fig 3B–E, Appendix Fig S3). The inhibitory effect of wild-type SOD1 or G93A mutant on miRNA biogenesis was consistent with this, but more limited (Fig 3F and G). The deleterious effect of higher-than-normal levels of
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**Figure 2. Different cellular stressors lead to impaired miRNA biogenesis.**

A Diagram depicting the working hypothesis. The DICER complex is composed of DICER, AGO2, PACT and TRBP, and the catalysis of pre-miRNA hairpins into mature ACO2-loaded miRNAs is schematically presented by a vertical arrow. Stress inhibits Dicing, resulting in the accumulation of substrate (pre-miRNA) and reduction in product levels (mature miRNA). The ratio of substrate to product, defined as “inhibition score,” approximates a value of 1 in the unmanipulated wild-type conditions. Inhibition score values greater than 1, reflect reduced DICER activity.

B–J Pre-miRNA (B, E, H) and miRNA (C, F, I) expression analysis and their corresponding inhibition score (D, G, J). NSC-34 cells treated with thapsigargin (10 nM for 24 h) or control carrier (DMSO) (B–D), paraquat (25 μM for 24 h) or control carrier (water) (E–F) and sodium arsenite (0.5 mM for 60 min) or control carrier (water) (H–I). Displayed are average and standard error of the mean (s.e.m.) for qPCR analyses of at least three independent experiments, normalized to the expression levels in control treatments. Pre-miRNA levels were normalized to beta-actin and Gapdh, and miRNAs were normalized to Snord70 and Snord70. P-values of qPCR were calculated via ANOVA statistics with DataAssist, and with two-sided Student’s t-test for inhibition score. Significant P-values are indicated by *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001.

Source data are available online for this figure.

wild-type FUS, TDP-43 or SOD1 is in accordance with the previous reports (Jaarsma et al., 2000; Wils et al., 2010; Igaz et al., 2011; Mitchell et al., 2013). Western blot analysis revealed that the levels of DICER, AGO2, PACT and TRBP did not change by the overexpression of wild-type, or mutant forms of FUS (Appendix Fig S4 related to Fig 3).
In order to test DICER activity directly, we devised a cell-free Dicing assay, using cell lysates and synthetic double-stranded RNA, carrying a quencher on a 25-nt passenger strand and a CY3 fluorophore conjugate on a 27-nt precursor guide strand. Dicing activity enables the dissociation of a fluorophore conjugate RNA from the quencher and fluorescence (Fig 4A). The assay demonstrated reduction in DICER catalytic activity in lysates of cells treated with sodium arsenite, paraquat or thapsigargin (Fig 4B and C). Importantly, overexpression of ALS-causing mutants or their respective wild-type forms also reduced DICER activity (Fig 4D). Noteworthy is that the in vitro assay probably provides an underestimate of the changes in DICER activity, due to the extensive dilution of soluble...

Figure 3. Over-expression of ALS-causing mutant proteins leads to impaired miRNA biogenesis.
A Diagram depicting the working hypothesis that expression of ALS-causing TDP-43, FUS or SOD1 attenuates DICER complex activity directly, or through induction of stress.
B-G Inhibition score of FUS R495X (B), wild-type FUS (C), TDP-43 A315T (D), wild-type TDP-43 (E), SOD1 G93A (F) or wild-type SOD1 (G). The calculated values are based on qPCR analysis results of pre-miRNAs, mature miRNAs of NSC-34 cell RNA, 72 h post-transfection, which is presented in Appendix Fig S3. The inhibition score of individual pre-miRNA:miRNA pairs was normalized to values in cells transfected with control vector. Shown are average and s.e.m. of > 3 independent experiments, except for pre-miRNAs in the study of FUS R495X (n = 2). P-values were calculated by two-sided Student’s t-test, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001. Source data are available online for this figure.
Figure 4. Stress or over-expression of ALS-causing mutants inhibits DICER activity in cell lysates.

A Diagram of the in vitro DICER activity assay, in which an annealed double-stranded RNA substrate is composed of a fluorophore-conjugated 27-nt guide strand and a 25-nt passenger strand conjugated to a quencher moiety. The DICER complex releases a 21-nt mature single-stranded guide RNA, whose fluorescence correlates with DICER activity.

B, C Dicing in vitro assay was performed in NSC-34 lysates after treatment with (B) sodium arsenite (0.5 mM for 60 min), paraquat (25 µM for 24 h) or control carrier (water), or alternatively (C) with thapsigargin (10 nM for 24 h) versus control carrier DMSO.

D Dicing in vitro assay was performed in HEK293 cell extracts 72 h post-transfection with the indicated plasmids.

E–G Inhibition score of DICER activity in cells transfected with ALS-causing mutants FUS R495X (E), TDP-43 A315T (F) or SOD1 G93A (G) and treated with carrier (buffer) or enoxacin (100 µM) for 72 h. The calculated values are based on qPCR analysis of pre-miRNAs, and mature miRNAs of NSC-34 cell RNA that is presented in Appendix Fig S5. Inhibition score of individual pre-miRNA:miRNA pairs was normalized to values in cells transfected with control vector.

Data information: (B–G) Shown are average and s.e.m. of > 3 independent experiments. P-values were calculated by two-sided Student’s t-test, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001.

Source data are available online for this figure.
DICER cofactors in the reaction buffers. Therefore, overexpression of ALS genes, mutation in ALS genes and the application of oxidative or ER stress all inhibit DICER catalytic activity.

Enoxacin ameliorates ALS-induced defects in pre-miRNA processing

Next, we hypothesized that enhancing Dicing complex activity might reverse the negative effect of the ALS-causing mutant proteins on miRNA processing. We tested this hypothesis by employing enoxacin, a fluoroquinolone antibiotic that is commonly used for the treatment of urinary tract and airway infections. Enoxacin is known to increase miRNA biogenesis via increasing the binding affinity of TRBP and pre-miRNAs (Shan et al., 2008; Melo et al., 2011). Intriguingly, the impairments in pre-miRNA processing that were evident in NSC-34 cells upon transfection with vectors expressing ALS-causing mutants FUS R495X, TDP-43 A315T or SOD1 G93A were partially ameliorated by applying enoxacin (100 μM for 72 h) to transfected cultures (Fig 4E–G and Appendix Fig S5). Therefore, enoxacin studies further support the view that DICER activity is impaired in culture models of ALS.

Stress granule formation controls miRNA biogenesis

To gain further mechanistic insight, we performed targeted mass spectrometry analysis of AGO2 and DICER protein interactions under cellular stress (Fig 5A). We examined a set of proteins that were co-immunoprecipitated with either AGO2 or DICER in a preliminary unbiased screen. Targeted mass spectrometry revealed enhanced interactions of AGO2 with DICER in cells that were treated with sodium arsenite, relative to basal conditions (Fig 5B and C). We report increased AGO2 binding to DICER, which was previously shown to decrease DICER complex activity (Tahbaz et al., 2004).

As DICER complex composition controls miRNA biogenesis (Lee & Doudna, 2012; Lee et al., 2013), we measured additional AGO2 and DICER interactions. HSP90 and co-chaperone p23, which regulate the interactions of DICER with AGO2 (Tahbaz et al., 2004; Pare et al., 2013), increased their interactions with DICER under stress (Fig 5B). These interactions potentially stabilize an intermediate DICER–AGO2–p23–HSP90 complex (Pare et al., 2013), thereby inhibiting DICER catalytic activity. In addition, cellular stress increased AGO2 and/or DICER protein interactions with SG components EIF2A, EIF3 (Kedersha et al., 2005) and EIF5A (Li et al., 2010) and with poly(rC)-binding proteins PCBP1 (hnRNP E1) and PCBP2 (hnRNP E2) (Li et al., 2012) (Fig 5B and C). PCBP1 and PCBP2 are iron chaperones (Shi et al., 2008; Nandakumar et al., 2011), found in SGs and P-bodies (Fujimura et al., 2008, 2009). PCBP2 binds to miRNA precursors and presents them to DICER for efficient processing (Li et al., 2012). Therefore, the binding of PCBP1 and 2 to AGO2 and DICER exposes an additional layer of complexity in the regulation of miRNA biogenesis under stress.

Finally, AGO2 is phosphorylated by the p38 mitogen-activated protein kinase on serine-378 under stress (Zeng et al., 2008), consistent with a 10-fold increase in AGO2 serine-378 phosphorylation in our system (Fig 5C). Intriguingly, p38 was reported to be persistently activated in ALS (Tortarolo et al., 2003). We further validated increased interactions of AGO2 with PCBP1, by AGO2 immunoprecipitation and Western blot analysis. DICER pull-down was marginally significant, and we could not validate the increased interaction of PACT with AGO2 (Fig 5D).

Taken together, these observations point toward substantial changes in the interactions of DICER and AGO2 with SG components. Next, we tested the hypothesis that SG formation may be sufficient to attenuate DICER complex activity. We overexpressed TIA1, a known initiator of stress granule formation (Kedersha et al., 1999), which reduced DICER catalytic activity in a cell-free assay (Fig 5E). EIF5A is a translation elongation factor, which is engaged with SGs under stress (Li et al., 2010) and also with DICER (Fig 5B). We demonstrated that EIF5A overexpression reduced DICER catalytic activity in a cell-free assay and in cells (Fig 5E and F and Appendix Fig S6A and B).

Several cellular stress-signaling cascades converge into the phosphorylation of Ser51 of the translation initiation factor 2 alpha (EIF2A), thereby driving SG formation (Hinnebusch, 2005; Kedersha et al., 2005). Because EIF2A interactions with AGO2 and/or DICER increased by stress (Fig 5B and C), we tested the implications of overexpressing a phosphomimetic version of EIF2A (S51D). Expression of EIF2A S51D resulted in inhibition of DICER activity (Fig 5G and Appendix Fig S6C and D), further linking stress signaling to reduced miRNA biogenesis. Accordingly, phosphorylation-resistant serine-to-alanine-mutated EIF2A (S51A) (Costa-Mattioli et al., 2007) enabled a partial recovery from thapsigargin-induced attenuation of DICER complex activity in mouse embryonic fibroblasts (Appendix Fig S7 related to Fig 5).

To test whether the experimental manipulations we used are sufficient to induce visible SGs, we utilized immunofluorescence microscopy. This study revealed the onset of TIA1-positive stress granules with a set of chemical SG inducers, or after transient transfection with vectors expressing wild-type or mutant forms of FUS, TDP-43 or SOD1. It may be that high plasmid copy number and the stress of the transient transfection contributed to the induction of SGs by wild-type FUS, which was not reported in a stably transfected cell line (Bosco et al., 2010; Dormann et al., 2010; Daigle et al., 2013).

In addition, overexpression of SG proteins that were sufficient to inhibit DICER activity, EIF5A, EIF2A S51D or TIA1 led to the formation of TIA1-positive SGs (Fig 6). Intriguingly, under all these conditions, AGO2 was co-localized with TIA1 in SGs, in accordance with the previous reports that observed AGO2 recruitment into SGs (Leung et al., 2006; Wu et al., 2011). Therefore, interactions of SG proteins with AGO2 and/or DICER directly contribute to the attenuation of DICER activity.

We next tested the effect of SGs on DICER activity via chemical induction or blockade of SG formation (Fig 7A). NSC-34 cells were treated with puromycin (1 μg/ml, 24 h), which drives dismantling of translating polysomes, thereby inducing SGs (Blobel & Sabatini, 1971). Puromycin decreased the efficacy of miRNA biogenesis in a cell-free Dicing assay (Fig 7B) and in cells, as measured by qPCR (Fig 7C and Appendix Fig S8A and B). Notably, puromycin (1 μg/ml, 24 h) reduced DICER and TRBP levels by 44 and 31%, respectively (Appendix Fig S9A and B related to Fig 7). Because the interpretation of the puromycin experiment can take into consideration reduction in DICER/TRBP levels, we sought a complementary approach to test whether inhibition of SG formation recovers miRNA biogenesis from ALS mutant-induced Dicing defects. Cycloheximide (CHX) is known to dissolve pre-formed SGs (Kedersha et al., 2000). The application of CHX to cultures transfected with

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Figure 5. Stress granule proteins interact with DICER complex components and modulate Dicing activity.

A Diagram depicting the working hypothesis that DICER complex interactions with stress granule proteins is modified by chemical stressors, overexpression of phosphomimetic EIF2A or overexpression of stress granule proteins.

B, C Targeted mass spectrometry analysis of proteins, co-immunoprecipitated with DICER-FLAG (B) or AGO2-FLAG (C), from HEK293 cells treated with sodium arsenite (0.5 mM, 60 min). Data are presented as ratio of averaged peptide counts from three different peptide standards per protein, relative to peptide levels in untreated cells. Note that two peptides were identified above threshold for EIF5A and for TRBP in the AGO2-IP, and AGO2 phospho-S387 is measured by a single peptide. Averages ± s.e.m.; *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 by two-sided Student's t-test, between arsenite and control treatment are denoted.

D Western blot study and densitometry of DICER, AGO2, PCBP1 and PACT after AGO2-FLAG immunoprecipitation, in three different biological replicates without or with sodium arsenite (0.5 mM, 60 min). Averages ± s.e.m., two-sided Student's t-test, ***P < 0.0001, for changes in AGO2 co-immunoprecipitated protein levels.

E In vitro Dicing activity assay in HEK293 cell lysate 72 h post-transfection with EIF5A or TIAR. Averages ± s.e.m.; ****P < 0.0001 by two-sided Student's t-test from > 3 independent biological replicates.

F, G Inhibition score of individual pre-miRNA:miRNA pairs of HEK293 cells transfected with EIF5A (F) or phosphomimetic form of EIF2A (S51D) (G). Calculated values are based on qPCR analysis of pre-miRNAs and mature miRNAs of NSC-34 cell RNA, presented in Appendix Fig S6. Inhibition score was normalized to values in cells transfected with control vector. Averages ± s.e.m.; *P < 0.05, **P < 0.01, ***P < 0.001 by two-sided Student's t-test from > 3 independent biological replicates.

Source data are available online for this figure.
### Figure 6. Localization of AGO2 to TIA1-positive stress granules.

**A** AGO2-FLAG HEK293 cells were treated with sodium arsenite (0.5 mM, 60 min), paraquat (25 μM, 24 h) or carrier (water), thapsigargin (10 nM, 24 h) or DMSO, puromycin (1 μg/ml and 2 μg/ml for 24 h) or carrier (water).

**B** Depicted are micrographs of AGO2-FLAG HEK293 cells transfected with plasmids as indicated and stained with antibodies for FLAG and TIA1, 72 h post-transfection.

**C** Micrographs of HEK293 cells transfected with plasmids as indicated and stained with antibodies for AGO2 and TIA1, 72 h post-transfection.

Data information: Scale bars indicate 10 μm.
FUS R495X-, TDP-43 A315T- or SOD1 G93A-expressing vectors were able to mitigate the reduction in DICER complex activity (Fig 7D–F and Appendix Fig S8C–H) and reduced the abundance of visible SG that were quantified by confocal microscopy (Fig 7G and H). Noteworthy, it is unlikely that the effect of CHX was due to a general inhibition of protein synthesis since puromycin, which equally inhibits translation, had an opposite effect (Fig 7B). Furthermore, with the CHX concentrations we used, we did not see a reduction in the ectopic expression of FLAG-TDP-43 A315T or GFP-FUS R495X and CHX was unable to affect miRNA levels when applied to untransfected NSC-34 cells (Appendix Fig S10A–C, related to Fig 7).

miRNAs down-regulation in the SOD1 G93A mouse model of ALS is reversed by enoxacin

We next extended our observations to the SOD1 G93A mouse model of ALS (B6SJL-Tg (SOD1*G93A)1Gur/J mouse strain) (Gurney et al., 1994). We detected down-regulation of several miRNAs in SOD1 G93A spinal cords at day 137, relative to the expression of the same miRNAs at day 77. These changes were not observed in age-matched and litter-matched wild-type controls (Appendix Fig S11A related to Fig 8). The mRNA and protein levels of AGO2, DICER, PACT and TRBP remained unchanged (Appendix Fig S11B and C related to Fig 8). The relatively mild reduction may be due to extracting RNA from the whole spinal cord tissue.

Recently, enoxacin was shown to increase the expression of miRNAs in the frontal cortex of rats with beneficial impact on depression (Smalheiser et al., 2014). Therefore, we tested whether miRNA levels are changed in vivo after oral application of enoxacin (n = 5) or carrier (water, n = 5) to SOD1 G93A male sibling-matched cohorts. Treatment starting on day 42 of the mouse life did not affect miRNA levels in wild-type motor cortices that were harvested at day 90, relative to controls. However, the levels of several miRNAs were up-regulated by enoxacin in SOD1 G93A motor cortices (Appendix Fig S11D and E related to Fig 8).

Enoxacin therapy is beneficial for neuromuscular function in the SOD1 G93A mouse model of ALS

We next tested whether enoxacin has a beneficial effect on neuromuscular function of SOD1 G93A mice (Gurney et al., 1994). We observed a ~7-day delay at the onset of neurological symptoms of the enoxacin-treated group (n = 40) relative to sibling-matched male cohort treated with carrier (water, n = 37) (Fig 8A). Consistently, weight peak and onset of weight decline, which is defined by the loss of 1 g bodyweight after the weight peak, were delayed in enoxacin-treated mice (Fig 8B and C). Kaplan–Meier survival analysis did not yield significant differences between the two groups (Fig 8D). However, evaluation of neurological status by employing a common neurological scoring system, with numerical values increasing from 0 to 4 as the disease progresses (Gill et al., 2009). Neurological scoring revealed that the enoxacin-treated cohort was superior to untreated controls (Fig 8E).

To further evaluate motor function, we performed automated quantitative gait analysis, using the CatWalk system (Neumann et al., 2009). Swing speed was significantly higher and stride length was significantly increased in the enoxacin-treated group, relative to untreated controls (ten matched siblings per group, Fig 8F and G). Additionally, the performance of the enoxacin-treated group was superior to untreated siblings in a rotarod test (n = 25 control, n = 29 enoxacin, Fig 8H). Therefore, enoxacin has a beneficial effect on multiple clinical parameters of the SOD1 G93A mouse model. Furthermore, even a lower enoxacin dose significantly improved gross strength on a hang-wire assay and the clinical neurological score, relative to controls (16 animals treated with enoxacin, 15 untreated controls; Fig 8I and J). Finally, enoxacin-treated animals (n = 6) moved significantly more than untreated controls (n = 5) in a fully automated infrared-based home cage locomotion assay (Fig 8K). Although enoxacin did not extend the lifespan of the SOD1 G93A model, an array of different assays revealed improved neuromuscular function, suggesting a beneficial profile. Noteworthy, riluzole, the only FDA-approved drug for ALS, lacks survival benefit in SOD1G93A mice (Scott et al., 2008; Li et al., 2013a). Therefore, as previously suggested (Scott et al., 2008), the predictive value of the aggressive SOD1 G93A model may be limited due to its 23 transgene copies in testing novel pharmacological interventions.

We also tested enoxacin in the TDP-43 A315T transgene (Wegorzewska et al., 2009). Enoxacin-treated TDP-43 A315T females displayed an improved neurological score, relative to a control cohort (nine treated animals, eight untreated controls; treatment started on day 42; Fig 8L), performed better on the Hang-wire test (Fig 8M) and displayed improved stride length (Fig 8N). Of note, characterization of the TDP-43 A315T mice was performed until the onset of clinical gastro-intestinal pathology, which leads to early death of this TDP-43 A315T mouse line and precludes the full development of ALS (Esmaeili et al., 2013).

Discussion

miRNAs function at an intriguing interface of cellular stress and disease (Leung & Sharp, 2007; Mendell & Olson, 2012; Emde & Hornstein, 2014). Accordingly, facets of miRNA malfunction in ALS or frontotemporal dementia were reported in the past: We demonstrated that loss of DICER is sufficient to cause progressive degeneration of spinal motor neurons (Haramati et al., 2010). In addition, specific miRNA genes are involved in feedback loops upstream or downstream of FUS (Morlando et al., 2012; Dini Modigliani et al., 2014) or TDP-43 (Buratti et al., 2010; Kawahara & Mieda-Sato, 2012). miRNAs were suggested to regulate neuromuscular junction repair (Williams et al., 2009; Valdez et al., 2014), to plausibly regulate neuro-inflammation in ALS (Koval et al., 2013) and may serve as circulating biomarkers (De Felice et al., 2014; reviewed in Droppelmann et al., 2014; Gascon & Gao, 2014).

The current work provides novel evidence that global down-regulation of miRNAs is a molecular commonality for genetically unrelated forms of human ALS and suggests that similar previous observations (Campos-Melo et al., 2013) are primarily due to changes in miRNA expression in motor neurons. The observations in human samples led us to investigate the mechanism. We demonstrated that decreased DICER catalytic activity is reducing miRNA levels. Several lines of evidence suggest that DICER complex activity is inhibited by the emergence of SGs: (i) The interaction of several RNA-binding proteins that are normally
Figure 7. Stress granule formation impacts on Dicing complex efficacy.

A) Diagram depicting the working hypothesis that DICER complex activity is attenuated by stress granules that can be initiated by polysome disassembly. Inhibition of DICER activity by overexpression of ALS-causing proteins can be mitigated by inhibiting stress granule formation with cycloheximide or by enhancing DICER activity with enoxacin.

B) Dicing in vitro assay was performed on lysates of NSC-34 cells, treated with puromycin (1 µg/ml for 24 h) or carrier (water). Average values of > 4 independent biological repeats; error bars represent s.e.m.; ***P < 0.001 by two-sided Student's t-test.

C–F) Inhibition score of individual pre-miRNA:miRNA pairs in NSC-34 cells treated with puromycin (1 µg/ml for 24 h) or carrier (water) (C) or cells transfected with ALS-causing FUS R495X (D), TDP-43 A315T (E) or SOD1 G93A (F) or control vector and treated with low-dose cycloheximide (CHX) (0.02 µg/ml) or carrier (water), starting at 6 h post-transfection. Cells were harvested 72 h post-transfection. Calculated values are based on qPCR analysis of pre-miRNAs and mature miRNAs presented in Appendix Fig S8. Inhibition score was normalized to values in cells transfected with control vector. Averages ± s.e.m.; *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 by two-sided Student's t-test from > 3 independent biological replicates.

G) AGO2-FLAG HEK293 cells transfected with FUS R495X or SOD1 G93A and HEK293 transfected with TDP-43 A315T vector versus control plasmid treated with cycloheximide (CHX, 0.02 µg/ml) or carrier and immunostained with antibodies as indicated 72 h post-transfection. Scale bars indicate 10 µm.

H) Percentage of HEK293 cells with stress granules overexpressing TDP-43 A315T or FUS R495X in the presence or absence of cycloheximide (0.02 µg/ml for 72 h). Average was calculated from > 100 cells in > 5 pictures per condition; error bars represent s.e.m.; ***P < 0.001 by two-sided Student's t-test.

Source data are available online for this figure.
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Figure 8. Beneficial impact of enoxacin on neuromuscular function of two different mouse ALS models.

A-H Oral application of enoxacin (800 mg/kg bodyweight/day; n = 40) or carrier (water; n = 37) to SOD1.G93A male mice started on day 42 of the mouse life (A) Onset of symptoms (neurological score 3); log-rank Mantel–Cox test, **P = 0.0002. (B) Weight peak, log-rank Mantel–Cox test, **P = 0.0009. (D) Kaplan–Meier survival plot reveals comparable life span of SOD1.G93A mice, regardless of therapy. Not significant (n.s.), by log-rank Mantel–Cox test. (F) Average neurological score, which increases with disease progression, per cohort (n = 40 enoxacin-treated, n = 37 controls), two-way ANOVA test, ***P < 0.0001. (control versus enoxacin). Post hoc Holm–Sidak tests did not reveal significant differences comparing control versus enoxacin at single time points. (F, G) Fully automated gait CatWalk analysis with ten matched siblings per group. (F) Four-paw swing speed on day 73, two-way ANOVA *P = 0.027 (control versus enoxacin). (G) Four-paw stride length on day 80, two-way ANOVA *P = 0.012 (control versus enoxacin). Post hoc Holm–Sidak tests did not reveal significant differences when comparing single-paw behavior. (H) Multiple time point rotarod performance, normalized to initial performance at day 92 of each individual (n = 25 control, n = 29 enoxacin). Two-way ANOVA with repeated measures for each individual, *P = 0.047 (control versus enoxacin), followed by post hoc Holm–Sidak tests, *P < 0.05, ***P < 0.001, ****P < 0.0001.

I-K Oral application of enoxacin 200 mg/kg bodyweight/day or carrier (water) to SOD1.G93A male mice started day 42 of the mouse life. (I) Hang-wire assay (n = 15 control, n = 16 enoxacin), two-way ANOVA (control versus enoxacin) *P = 0.0027, followed by post hoc Holm–Sidak tests of different time points within groups, **P < 0.001, ***P < 0.0001. Lower hang-wire score indicate greater strength. (J) Neurological score, which increases with disease progression (n = 15 control, n = 16 enoxacin), two-way ANOVA (control versus enoxacin) *P = 0.0084, followed by post hoc Holm–Sidak test reveals a significant difference at 20 weeks of age **P < 0.01. (K) Fully automated infrared-based home cage locomotion analysis (InfraMot, TSE-Systems) at 30-min intervals over a period of 46 h, days 129–130 (n = 5 control, n = 6 enoxacin). Two-way ANOVA (control versus enoxacin), ***P = 0.0004.

Data information: All error bars, s.e.m. Note that in (A–C, D), x-axis starts at days 60 and 100, respectively. Discontinuation of y-axis in (F, G, N) is marked by oblique crossing line.

Source data are available online for this figure.
found in stress granules with DICER or AGO2 increases upon stress. (ii) Initiating SGs by overexpressing EIF5A or TIAR, the phospho-
mimetic form EIF2A, or by puromycin, was sufficient to reduce DICER activity in cells and in cell lysates. (iii) AGO2, a cofactor of DICER, co-localizes with TIA1 in cytoplasmic SGs after transfection of ALS-causing genes. (iv) Blocking SGs with cycloheximide or with non-phosphorylatable serine-to-alanine S1 mutant of EIF2A reversed the negative effect of ALS-causing mutants on miRNA biogenesis. Puromycin and cycloheximide are two inhibitors of translation, yet they display opposite effects both on pre-miRNA processing and, respectively, on SG formation. Thus, the protective activity of cycloheximide in motor neurons (Yang et al, 2013) is consistent with its ability to inhibit SG formation and to improve miRNA biogenesis. (v) These observations are in line with the previous identification of TDP-43 and FUS in SGs and with the fact that TDP-43 binds DICER and AGO2 directly (Kawahara & Mieda-Sato, 2012), providing a conceivable explanation also for the stronger molecular phenotypes observed with TDP-43 and FUS, relative to SOD1.

Therefore, we suggest that stress and SG formation initiate dynamic changes in DICER interactions with its co-factors, in a way that diminishes DICER complex activity. This may be because of acquired or lost interactions under stress and may be related to AGO2 post-translational modifications (Leung et al, 2006; Qi et al, 2008; Zeng et al, 2008; Shen et al, 2013) or to increased AGO2 binding to DICER, which counter-intuitively decreases DICER activity (Tabbazi et al, 2004).

Enoxacin increases the binding affinity of TRBP to pre-miRNAs, thereby augmenting pre-miRNA processing (Shan et al, 2008; Melo et al, 2011). We demonstrated that impaired miRNA biogenesis is mitigated by enoxacin with beneficial clinical outcome on the neuromuscular performance of two different ALS mouse models. We did not observe any lifespan extension, but also Riluzole, the only FDA-approved drug for ALS, lacks survival benefit in SOD1 G93A mice (Scott et al, 2008; Li et al, 2013a). Therefore, as previously suggested (Scott et al, 2008), the predictive value of the aggressive SOD1 G93A model may be limited in testing novel pharmacological interventions. Furthermore, SOD1 effect on DICER activity may be more limited than the effects mediated by FUS or TDP-43, which are RNA-binding proteins that are directly engaged in SGs.

Nonetheless, if DICER and miRNAs are indeed instrumental in the pathogenesis of ALS in humans, as we suggest, these observations hold promise for future intervention by validating DICER as a new therapeutic target. Furthermore, enoxacin is a particularly intriguing candidate for clinical assessment, based on its established safety and pharmacokinetic profile and because several forms of ALS converge in the down-regulation of miRNAs. However, more effective DICER agonists might exhibit more dramatic beneficial effects in ALS in the future.

miRNAs impart robustness to cellular programs in development and in the adult life (Hornstein & Shomron, 2006; Pelaez & Carthew, 2012; Cassidy et al, 2013; Emde & Hornstein, 2014). Accordingly, even moderate DICER and miRNA insufficiency contributes to failed homeostatic mechanisms, allowing the activation of myriad aberrant pathways that were otherwise repressed under normal miRNA activity. The consequences may contribute particularly to diminished stress resistance, creating a vicious cycle, which further increases neuronal stress up to the threshold of malfunction and death. Accordingly, in one example, even mild but long-term down-regulation of brain-enriched miR-124 affected AMPA expression and frontotemporal dementia phenotype (Gascon et al, 2014).

The sensitivity of DICER activity to cellular stress is not limited to motor neurons and may be probably relevant to other tissues under chronic stress, where decrease in miRNA or DICER abundance was reported (Kaneko et al, 2011; Mori et al, 2012; Inukai & Slack, 2013; Nidadavolu et al, 2013). Thus, evaluating whether stress impinges on DICER activity in other disease states will be valuable. The variability in miRNA biogenesis with different stressors or by using different cell types suggests additional factors that are currently unknown.

In summary, we provide evidence for the control of miRNA biogenesis at the level of DICER complex activity, by several stress-related cellular changes. The consecutive reduction in miRNA levels unifies various forms of ALS, suggesting that the development of novel therapies might focus on DICER or miRNAs as common critical effectors of neuronal integrity.

Materials and Methods

Human tissue miRNA analysis

Human nervous tissues were acquired by way of an Investigational Review Board and Health Insurance Portability and Accountability Act compliant process. Sporadic ALS nervous systems were from patients who had met El Escorial criteria for definite ALS. Informed consent was obtained from all subjects. Tissue measurements were carried out post-mortem and conform to the WMA Declaration of Helsinki and the Department of Health and Human Services Belmont Report. Sporadic ALS nervous systems were from patients who had bulbar or arm-onset disease and caudal progression, and thus, the lumbar regions had relatively abundant residual motor neurons. Tissue collections were as previously described: Completed within 4–6 h of death, the motor system was dissected, embedded in OCT and stored at –80°C. RNA quality was assessed using microelectrophoresis on an Agilent 2100 Bioanalyzer as previously described. Microdissected cell punches were captured on CapSure™ Macro LCM Caps (Arcturus Bioscience). 35 to 50 cryosections were cut at a thickness of 9 µm in a –18°C cryotome and placed onto uncharged glass slides. The sections were returned to –80°C for a minimum of 3 h. They were stained with cresyl violet acetate in a 10-step, timed, nuclease-free immersion process. Motor neurons were microdissected using a Pixcell IIe Laser Capture Microdissection (LCM) System (Arcturus Bioscience) and CapSureTM Macro LCM Caps (Applied Biosystems). Each LCM session collected 500–1,000 motor neuron punches and lasted < 2.5 h. Total RNA was isolated using the RNAqueous Micro kit (Life Technologies) as per the manufacturer’s procedure. RNA quality was assessed by spectrophotometric analysis using a NanoDrop ND-1000 spectrophotometer and an Agilent 2100 Bioanalyzer. Expression of 667 microRNAs was assessed using the microRNA TaqMan® qPCR Megaplex pools array on an ABI 7900HT Fast Real-Time PCR System (Life Technologies). Technical replicates were run after pre-amplification using the manufacturer’s protocol. Relative quantification was calculated by comparing arithmetic mean of three small RNA control genes nuclear RNU48/SNORD48,
RNU44/SNORD44 and splicesome-related U6. Two-way ANOVA test was performed with DataAssist Software (Life technologies Inc.). In situ hybridization was performed on 7-µm sections of frozen spinal tissue from lumbar regions, briefly rinsed in 4% paraformaldehyde and incubated in 2 µg/ml proteinase K. Slides were then incubated for 10 min in 0.13 M 1-methylimidazole (Sigma), 300 mM NaCl, pH 8.0 and further fixed in EDC (Sigma) following Pena et al (2009), acetylated for 30 min in a solution of freshly prepared 0.1 M triethanolamine and 0.5% (v/v) acetic anhydride. Hybridization of sections with 4 pmol of 5’ DIG-labeled miR-9 and miR-124 LNA probes followed manufacturer’s instructions (Exiqon).

**Tissue culture vectors and small molecules**

HEK293 cells (American Type Culture Collection) and NSC-34 cells (Cellutions Biosystems Inc.) were cultured in high-glucose DMEM (Gibco), 1% l-glutamine, 10% fetal calf serum and 1% penicillin/streptavadin. Mouse embryonic fibroblasts (MEFs) were cultured in DMEM (Gibco), 15% fetal calf serum, l-glutamine, 1% penicillin/streptavadin and sodium pyruvate.AGO2-FLAG-expressing HEK293 cells were a gift of Markus Landthaler. EIF2A S51A mouse embryonic fibroblasts were provided by Prof. Chaim Kahana, Weizmann Institute of Science, Israel. Mouse embryonic motor neurons were harvested as described in Milligan and Gifondorwa (2011). Expression vectors are as follows: EIF2A 3’ (S51D) from David Ron (Addgene plasmid # 21809); pBabe_puro_DEST_Flag_EIF5A2 from William Hahn (Addgene plasmid # 45266); and pLMM60-Tiar from David Sabatini (Addgene plasmid # 38244). Wild-type and R495X FUS vector were provided by Daryl Bosco (University of Massachusetts, Boston), wild-type and A315T TDP-43 vectors by Markus Landthaler (MDC Berlin, Germany), and wild-type and G93A SOD1 vectors by Yoram Groner (WIS, Israel). DICER-FLAG and AGO2-FLAG plasmids were kindly provided by Narry Kim (Seoul National University, South Korea). Enoxacin was supplied by Sigma-Aldrich (E3764-5G) or Buckton-Scott (Germany). Thapsigargin, pararaut, cycloheximide and puromycin are purchased from Sigma-Aldrich, 20-500-120). Primary antibodies used are as follows: anti-TDP-43 (10782-2-AP, Proteintech) 1:1,000; TLS/FUS (cat. 611385, BD Transduction Laboratories) 1:500; and anti-eIF2α (sc-11386, Santa Cruz Biotechnology) 1:200; anti-TRBP (ab42018) 1:1,000; anti-PCBP1 (ab75749) 1:1,000; anti-GRAPDH (Ambion, AM4300) 1:4,000; anti-phospho-eIF2α (Ser51) (Cat# 04-342EMD, Millipore) 1:500; anti-eIF2α (sc-11386, Santa Cruz Biotechnology) 1:200; anti-TLS/FUS (cat. 611385, BD Transduction Laboratories) 1:500; and anti-TDP-43 (10782-2-AP, Proteintech) 1:1,000.

**RNA analysis**

RNA was extracted using Tri reagent (Molecular Research Center Inc.) or Qiazol with consecutive use of the miRNeasy Mini Kit (Qia-gen). Pre-miRNA and miRNA reverse transcription was performed using miScript Kits No. I & II (Qiagen). Quantitative analysis of miRNA and pre-miRNA expression was performed in > 3 independent biological repeats and additionally in technical duplicates with StepOnePlus quantitative Real-Time PCR System, and two-way ANOVA test was performed with DataAssist Software (Life technologies Inc.). Primers are described in Table E1V and additional pre-miRNA sequences are from Jiang et al (2005).

**Cell-free dicing assay**

Dicing assay is described in Melo and Melo (2014) with the following modifications: dsRNA substrate for measuring DICER activity was annealed from 5’ Phos/mCmU rCmArU mUrUUrC rCrUUrG mGrUmA rUmGrA mCrArA rCrGrA mAmU/3’IA8kBkFQ and 5’Cy3/ mAmU mUrCrG rUrUGr mUrCmA rUmArC rCrArA rArUmG rAmGmC rU (Integrated DNA Technologies). Cells were lysed via sonication in hypotonic solution, with consecutive DICER assay reaction assembly by mixing 3 µl of 10× Buffer K (200 mM Heps–KOH, pH 7.0, 20 mM dithiothreitol (DTT), 20 mM MgCl2; 3 µl of 10 mM ATP; 1.5 µl of 0.5 M, freshly dissolved, creatine phosphate; 0.9 µl of 1 mg/ml creatine phosphate kinase; 0.6 µl of ribonuclease inhibitor (40 units/µl); 3 µl of labeled dsRNA (50 nM); and 18 µl of cell extract. DICER assay mix was incubated at 30°C for 2 h and diluted with 100 µl of ultra-pure water, and then, 10 µl was transferred to 384-multiwell plate (10 technical repeats for each reaction) and measured on PHERAstar FS microplate reader (BMG Labtech). Pre-anaeelled ssRNA Cy3 fluorescent oligo served as positive control for calibration curve. Annealed and quenched dsRNA without cell extract served as negative control for quantification of background fluorescence.

**Western blot analysis**

Cell extracts or immune-purified proteins were eluted by boiling in X5 sample buffer (60 mM Tris–HCl pH 6.8, 25% glycerol, 2% SDS, 14.4 mM β-mercaptoethanol, 0.1% bromophenol blue) for 5 min. The samples were separated via 8% SDS–polyacrylamide gel electrophoresis at 120 V for 70 min, electro-transferred onto nitrocellulose membrane (Whatmann, 10401383) at 100 V for 1 h, stained with Ponceau (Sigma, P7170) to asses transfer quality and then blocked for 1 h at room temperature with 5% milk protein in PBS + 0.05% Tween-20 (0.05% PBST). Blocked membranes were then incubated with shaking at 4°C overnight with primary antibodies in 5% bovine serum albumin, 0.02% sodium azide and five drops of phenol red in 0.05% PBST. Membranes were next washed 3 times for 5 min at room temperature with 0.05% PBST and then incubated with horseradish peroxidase-conjugated species-specific secondary antibodies in 5% milk for 1 h at room temperature. Membranes were washed three times for 5 min in 0.05% PBST and visualized by ImageQuant™ LAS 4000 (GE Healthcare Life Sciences) using EZ-ECL Chemiluminescence detection kit (Biological Industries, 20-500-120). Primary antibodies used are as follows: anti-DICER (ab13502) 1:500; anti-AGO2 (ab32381) 1:500; anti-PACT (ab87479) 1:1,000; anti-TRBP (ab42018) 1:1,000; anti-PCBP1 (ab74793) 1:1,000 from Abcam; anti-GAPDH (Ambion, AM4300) 1:4,000; anti-phospho-eIF2α (Ser51) (Cat# 04-342EMD, Millipore) 1:500; anti-eIF2α (sc-11386, Santa Cruz Biotechnology) 1:200; anti-TLS/FUS (cat. 611385, BD Transduction Laboratories) 1:500; and anti-TDP-43 (10782-2-AP, Proteintech) 1:1,000.

**Mass spectrometry**

For mass spectrometry, HEK293 cells that stably express AGO2-FLAG (n = 3 for each group) or DICER-FLAG-transfected cells (48 h post-transfection, n = 4 for each group) were untreated or stressed with 0.5 mM sodium arsenite for 60 min before harvest. IP was performed with the FLAG-IP1 Kit (Sigma-Aldrich). Negative controls were HEK293 cells not containing a FLAG construct. Eluted proteins and the FLAG peptide were filtered using 3-kDa molecular weight cutoff spin columns (Amicon, Millipore). Buffer was exchanged to 50 mM ammonium bicarbonate (Sigma-Aldrich) in the same step. Proteins were reduced by addition of dithiothreitol (Sigma-Aldrich) to a final concentration of 5 mM and incubation for 30 min at 60°C and
alkylated with 10 mM iodoacetamide (Sigma-Aldrich) in the dark for 30 min at 21°C. The proteins were then digested using trypsin (Promega; Madison, WI, USA) at a ratio of 1:50 (w/w trypsin/protein) for 16 h at 37°C. Digestion was stopped by addition of 1% trifluoroacetic acid (TFA). Samples were stored in −80°C until analysis. Each sample was spiked with the mixture of 51 heavy isotopically labeled peptides (JPT Technologies). Heavy labels included U-13C6; U-15N4 for peptides terminating with Arg and U-13C3; U-15N2 for Lys. ULC/MS grade solvents were used for all chromatographic steps. Each sample was loaded using split-less nano-ultra-performance liquid chromatography (10k psi nanoAcquity; Waters, Milford, MA, USA). Mobile phase was as follows: A) H2O + 0.1% formic acid and B) acetonitrile + 0.1% formic acid. Desalting of samples was performed online using a reverse-phase C18 trapping column (180 μm i.d., 20 mm length, 5 μm particle size; Waters). The peptides in samples were separated using a C18 T3 HSS nano-column (75 μm i.d., 250 mm length, 1.8 μm particle size; Waters) at 0.3 μl/min. Peptides were eluted from the column and into the mass spectrometer using the following gradient: 3–30% B in 90 min, 30–35% in 10 min, 35–90% B in 5 min, maintained at 90% for 5 min and then back to initial conditions. The analytical column was coupled with a quadrupole orbitrap mass spectrometer (Q Exactive, Thermo Scientific), via a nano-ESI interface. Parallel reaction monitoring of differential peptide expression. Peptide sequences for mass spectrometry are described in Table EV1.

**Immunocytofluorescence microscopy**

Wild-type or stably transfected AGO2-FLAG HEK293 cells were cultured on poly-A-lysine-coated coverslips and further subjected to transfection or treated with stress-inducing chemicals. For microscopy, cells on coverslips were rinsed with PBS, fixed with 4% formaldehyde, permeabilized with 0.1% Triton X-100 (Sigma-Aldrich, St. Louis, MO), blocked with PBS, 2% BSA and incubated with anti-TIA1 antibody (sc-1757, Santa Cruz Biotechnology), anti-Flag antibody (F1804, Sigma-Aldrich) and anti-AGO2 (ab32381, Abcam) overnight. Coverslips were washed three times with PBS and incubated with secondary antibodies (Cy2 donkey anti-goat 705-995-147 and Cy5 donkey anti-mouse 715-175-150, Jackson Immunoresearch) for 1 h. Coverslips were washed 3 times with PBS, mounted with Fluoroshield™ with DAPI (F6057, Sigma) and captured with a LSM 710 confocal microscope (Carl Zeiss AG).

**In vivo enoxacin study**

All experiments were performed according to Weizmann Institute of Science guidelines and IACUC approval (protocol No. 03040512-1). Mice, B6SJL-Tg(SOD1*G93A)1Gur/J, which were first reported in Gurney et al. (1994), carry a hemizygous high transgene copy number of the SOD1 G93A mutant on a B6SJL hybrid background (Jackson Laboratories, 002726). The TDP-43 A315T mouse line (Wegorzewska et al., 2009) was on a C57BL/6J background (Jackson Laboratories, 010700). All experiments were performed sibling-matched, except the mice in the 200 mg/kg bodyweight SOD1 G93A enoxacin group, in which mice were randomly assigned to treatment groups. Animals were group-housed on a 12-h light–dark cycle. Food and water were provided ad libitum. Food pellets at cage floor, long sipper tubes on bottles and nutrient gel were used to ease accessibility to nutrition when mice deteriorated according to Gill et al. (2009). RotaRod™ performance (San Diego Instruments) was evaluated at ramp speed 4, 40 rpm. Data are presented as the average latency to fall off the rotarod of three independent rounds per time point and are normalized to average initial performance at pre-symptomatic stage for each individual for each time point. A fully automated gait analysis with the CatWalk XT™ (Noldus) system was performed. Data were obtained of > 4 independent rounds on the apparatus. For hang-wire muscle strength assessment, mice were allowed forelimb grip onto a 2-mm-thick horizontal metal wire, suspended 80 cm above surface. Ability to successfully raise hindlimbs to grip the wire and crawling to the end of the wire was scored: 1 = successful four-paw grip and crawling to the end of the wire within < 60 s; 2 = four-paw grip for > 15 s without crawling; 3 = failure to successfully establish hindlimb grip in 60 s; and 4 = inability to sustain forelimb grip for 60 s. Each mouse was evaluated in two sessions on two consecutive days, consisting of three trials each, with a 15-min inter-trial interval. The second day was used to obtain the data as displayed. Neurological score was assessed according to Gill et al. (2009): 0 = > 2 s extension of hindlimbs away from lateral midline when mouse is suspended by its tail 2–3 times; 1 = collapse or partial leg collapse toward midline or hind leg trembling during tail suspension; 2 = toes curl under at least twice during a 12-inch walk, or any part of foot is dragging along cage/table bottom; 3 = rigid paralysis or minimal joint movement, foot not being used for forward motion; and 4 = humane endpoint, mouse cannot right itself within 30 s on either side. The locomotion of animals was quantified over a period of 46 h in the home cage, by automated sensing of body-heat image using an InfraronMot (TSE-Systems). Individual animal movements were summed up every 30 min. Animal statistics: rotarod performance was evaluated using the two-way ANOVA with repeated measures for each individual and post hoc Sidak–Holm tests. Weight peak, weight decline, onset of symptoms and survival were calculated with log-rank (Mantel–Cox) test. All other tests employed two-way ANOVA with post hoc Sidak–Holm tests. All animal statistics were calculated with GraphPad Prism 6 statistics program.

**Expanded View**

For this article is available online: http://emboj.embopress.org

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Author contributions
AE conceived and led the research, performed all non-human experimental activities and data analysis, including molecular biology and animal studies. RTL and JMR provided human autopsy material. L-LL, ASi and BA performed laser capture microdissection and human tissue RNA analysis. NR performed microscopic studies, developed in situ hybridization study. ASa and YL performed mass spectrometry. IR, JA, IM, RS, IZB-D and SMH helped conducting research or provided critical input for scientific interpretations. JMR, TM and EH conceived research and supervised the study. AE and EH developed the interpretations presented and wrote the manuscript with comments from JMR and TM. JMR is the corresponding author for the human analysis, and EH is the corresponding author for all other facets of the study. TM and EH share senior authorship.

Conflict of interest
The authors declare that they have no conflict of interest.

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