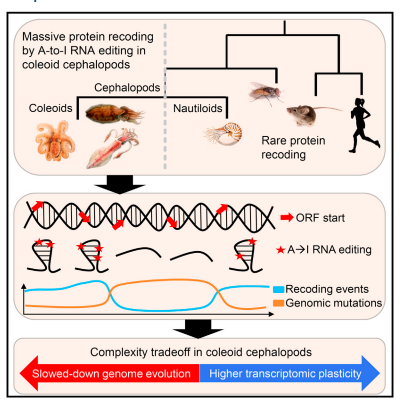


Trade-off between Transcriptome Plasticity and **Genome Evolution in Cephalopods**

Graphical Abstract



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In Brief

Behaviorally complex cephalopods use extensive RNA editing to diversify their neural proteome at the cost of limiting genomic sequence flexibility and evolution.

Highlights

- Unlike other taxa, cephalopods diversify their proteomes extensively by RNA editing
- Extensive recoding is specific to the behaviorally complex coleiods
- Unlike mammals, cephalopod recoding is evolutionarily conserved and often adaptive
- Transcriptome diversification comes at the expense of slowed-down genome evolution





Trade-off between Transcriptome Plasticity and Genome Evolution in Cephalopods

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SUMMARY

RNA editing, a post-transcriptional process, allows the diversification of proteomes beyond the genomic blueprint; however it is infrequently used among animals for this purpose. Recent reports suggesting increased levels of RNA editing in squids thus raise the question of the nature and effects of these events. We here show that RNA editing is particularly common in behaviorally sophisticated coleoid cephalopods, with tens of thousands of evolutionarily conserved sites. Editing is enriched in the nervous system, affecting molecules pertinent for excitability and neuronal morphology. The genomic sequence flanking editing sites is highly conserved, suggesting that the process confers a selective advantage. Due to the large number of sites, the surrounding conservation greatly reduces the number of mutations and genomic polymorphisms in protein-coding regions. This trade-off between genome evolution and transcriptome plasticity highlights the importance of RNA recoding as a strategy for diversifying proteins, particularly those associated with neural function.

INTRODUCTION

It is generally assumed that genetic information passes faithfully from DNA to RNA to proteins. Proteome complexity, however, depends on a diverse set of post-transcriptional processes that modify and enrich genetic information beyond the genomic blueprint. RNA editing is one such process. Adenosine deamination to inosine by the ADAR family of enzymes (adenosine deaminases acting on RNA) is the most common form of editing among animals (Bass, 2002; Nishikura, 2016). Because inosine is recognized as guanosine during translation (Basilio et al., 1962), this process has the capacity to recode codons and fine-tune protein function. However, it seldom does so. Transcriptome-wide screens have revealed that only \sim 3% of human messages and 1%-4% of those from Drosophila harbor a recoding site (Ramaswami and Li, 2014; St Laurent et al., 2013; Xu and Zhang, 2014; Yu et al., 2016). Even more surprising is the limited extent to which this process is conserved. There are only about 25 human transcripts that contain a recoding site that is conserved across mammals (Pinto et al., 2014), and only about 65 recoding sites conserved across the Drosophila lineage (Yu et al., 2016). In C. elegans, only a few putative recoding sites have been identified, some of which were not validated (Goldstein et al., 2017; Washburn et al., 2014; Whipple et al., 2015; Zhao et al., 2015). These data support the hypothesis that recoding by RNA editing is mostly neutral or detrimental and only rarely adaptive (Xu and Zhang, 2014).

Recently, we reported an apparent exception: squid contain an unusually high level of recoding, with the majority of mRNAs in the nervous system harboring at least one event (Alon et al., 2015). This intriguing but anecdotal result raised fundamental questions about the nature of recoding in these organisms. Does the massive RNA-level recoding translate into proteome diversification? Is it simply a neutral byproduct of a promiscuous ADAR tasked with another function or adaptive, providing a functional advantage? Finally, is it related to behavioral sophistication?

Cephalopods are diverse and can be divided into the behaviorally complex coleoids, consisting of squid, cuttlefish, and octopus, and the more primitive nautiloids. In this paper we show that in neural transcriptomes extensive A-to-I RNA editing is observed in the behaviorally complex coleoid cephalopods but not in nautilus. The edited transcripts are translated into protein isoforms with modified functional properties. By comparing editing across coleoid taxa, we found that, unlike the case for mammals, many sites are highly conserved across the lineage and undergo positive selection, resulting in a sizable slowdown of coleoid genome evolution.

RESULTS

Extensive Recoding Is an Invention of Coleoid Cephalopods

To assess the level of recoding via A-to-I RNA editing in cephalopods, we analyzed matching DNA and RNA samples of



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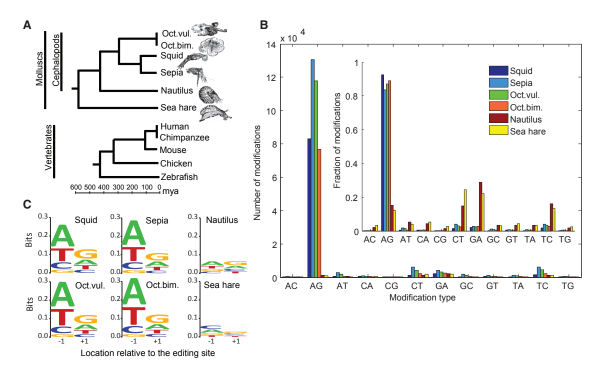


Figure 1. Extensive Recoding Is an Invention of Coleoid Cephalopods

(A) The species studied span the cephalopod evolutionary tree, as well as sea hare (Aplysia californica), as an outgroup (top). For comparison, a representative tree for vertebrates is shown (bottom), constructed based on divergence times estimated in Hedges et al. (2006).

(B) Tens of thousands of A-to-I editing sites (identified as A-to-G DNA-RNA mismatches) are detected in squid, sepia, and the two octopus species (see Tables S1-S4 for more details). The noise level (estimated by the number of G-to-A mismatches) is rather low. In contrast, in nautilus and sea hare, no enrichment of A-to-G mismatches is observed (inset).

(C) The nucleotides neighboring the detected editing sites show a clear pattern consistent with known ADAR preference (Alon et al., 2015; Eggington et al., 2011; Kleinberger and Eisenberg, 2010) for the extensively recoded coleoid species - squid, sepia, and the two octopus species - but not in nautilus or sea hare. The motif is characterized by under-representation of G upstream to the editing site (relative location -1) and over-representation of G in the downstream base (the height of the entire stack of letters represents the information content in bits; the relative height of each letter represents its frequency).

individual animals from species that span the cephalopod evolutionary tree. We studied four members of the coleoid cephalopod subclass (soft-bodied cephalopods): two octopuses (Octopus vulgaris and Octopus bimaculoides), a squid (Doryteuthis pealeii), and a cuttlefish (Sepia oficianalis), as well as a nautiloid (Nautilus pompilius) and a gastropod mollusk (Aplysia californica), as an evolutionary outgroup. Cephalopods emerged in the late Cambrian period, roughly at ~530 million years ago (mya), and the divergence of nautiloids from coleoides is estimated to have occurred at 350-480 mya (Kröger et al., 2011). The coleoides diverged to Vampyropoda (octopus lineage) and the Decabrachia (squid and cuttlefish lineage) at ~200-350 mya (Albertin et al., 2015; Kröger et al., 2011). Divergence of squid from Sepiida is estimated to have occurred at 120-220 mya (Checa et al., 2015). The two octopus species used in this study, Octopus vulgaris and Octopus bimaculoides, have been shown to be closely related using mitochondrial DNA and are in some cases even indistinguishable, depending on the geographical origins of the specimens (Söller et al., 2000). The divergence time between the gastropod species Aplysia californica and cephalopods is estimated to be 520-610 mya (Kröger et al., 2011). A general representation of the phylogenetic relations between the species is shown in Figure 1A.

A full genome sequence is not available for the cephalopod species used in this study (except for Octopus bimaculoides; see below). Thus, to detect editing sites, we used a genomeindependent method (Alon et al., 2015) that focuses specifically on the coding regions of the transcriptome. Briefly, RNA-seq data (174-366 million reads per species; Table S1, also see Alon et al., 2015) was utilized to assemble a de novo transcriptome (Grabherr et al., 2013), and the coding sequences were identified by comparison with Swiss-Prot (Bairoch et al., 2005) open reading frames (Table S2). RNA and DNA reads were then aligned to the assembled transcriptome (using Bowtie2 [Langmead and Salzberg, 2012] with local alignment configuration and default parameters). To detect editing events, we looked for systematic mismatches between RNA and DNA reads within the coding part of the transcriptome, filtering out those that stem from sequencing errors or genomic polymorphisms (see STAR Methods for more details). The A-to-G DNA-to-RNA mismatches that are identified by this process could result from A-to-I RNA editing, whereas other types of mismatches provide an estimate of our false-detection rate.

For sepia, squid, and the two octopus species, most mismatches (>80%) detected by the above approach were A-to-G mismatches, and the noise level, estimated by the number of

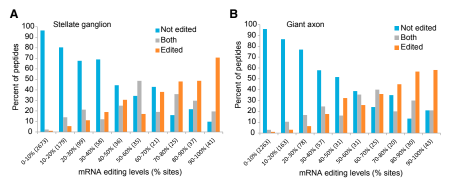


Figure 2. Proteomic Validation of Recoding by RNA Editing

We analyzed pentides identified by mass-spectrometry analysis of two squid tissues, looking for evidence of recoding. For each site covered by one or more peptides, we marked whether the edited, non-edited, or both versions of the peptide are observed. The distribution is presented, binned by the predicted RNA-editing level (as measured from RNA-seq data). In parentheses are the numbers of recoding sites analyzed in each editing-level bin. The proteomic recoding level follows closely the predicted RNA-editing level. Altogether, this experiment validated protein recoding in 432 sites in two tissues:

(A) Squid stellate ganglion, where 320 of the 3,204 single-site peptides (10.0%) were shown to be edited. (B) Squid giant axon (giant fiber lobe), where 283 of the 2,741 single-site peptides (10.3%) were shown to be edited.

G-to-A mismatches, is rather low at 2%-3% (Figure 1B). Furthermore, the residues surrounding the detected A-to-G sites exhibit a sequence pattern consistent with the known preferences for ADARs (Eggington et al., 2011; Kleinberger and Eisenberg, 2010) (Figure 1C). We thus attribute these mismatches to A-to-I RNA-editing events and obtain 80-130 thousand editing sites in protein-coding regions (Tables S3 and S4). Remarkably, results from nautilus and Aplysia are in sharp contrast. First, we found only 1,150 and 933 A-to-G mismatches for these species, much less than for the octopus, squid, and sepia. Moreover, there is no excess of A-to-G mismatches over other events (Figure 1B; Tables S3 and S4), and the residues surrounding the detected A-to-G sites do not exhibit any sequence preference (Figure 1C). Thus, the A-to-G mismatches found in nautilus and Aplysia are likely to be (mostly) noise, with very few, if any, editing sites. Accordingly, editing within the coding sequence of these species is orders of magnitudes lower than for the octopus, squid, and sepia. These data suggest that extensive recoding through RNA editing evolved along the coleoid lineage. As all of the cephalopod groups that separate coleoids and nautiloids are now extinct (e.g., belemnites and ammonoids), it will be difficult to pinpoint a more exact time for the emergence of extensive RNA editing.

Proteomic Validation of Recoding Sites

Sanger-sequencing validation of the sites detected by the present scheme was previously reported (Alon et al., 2015). Here, we employed mass-spectrometry analysis to further test whether the multitude of novel RNA isoforms created by extensive RNA editing are translated into proteins, resulting in extensive proteome diversification by recoding. We analyzed squid giant axon and stellate ganglion samples, looking for peptides translated from RNA that include editing sites. To simplify the analysis, we considered only peptides that include a single non-synonymous (recoding) editing site and checked whether the edited, non-edited, or both versions of the peptide were observed. For squid stellate ganglion, a total of 74,146 unique peptides were detected, 4,115 of which harbor 5,617 recoding sites, and 3,204 peptides included a single predicted site. Of these, 320 sites (10.0%) were shown to be edited (174 cases where both the pre-edited and edited versions are observed, and 146 found only in the edited version), including most of the sites predicted to be edited at high levels. Similarly, for squid giant axon 58,403 unique peptides were detected, 3,579 of which harbor 4,956 predicted recoding sites, and 2,741 peptides included a single predicted site. Of these, 283 sites (10.3%) were shown to be edited (160 cases where both pre-edited and edited versions are observed, and 123 found only in the edited version). Altogether, this experiment validated 432 protein-recoding sites. The fraction of sites validated correlated very well with the editing level predicted from RNA-seg data (Figure 2).

Note that the shotgun proteomics method used here provides only partial coverage of the tryptic peptides generated by the proteolysis (Michalski et al., 2011). This is demonstrated by the fact that ~90% of the recoded amino acids are completely missing from our data, regardless of their editing state. Accordingly, lack of peptide evidence for an edited or unedited form of a given site cannot be considered as evidence for this isoform not being present. However, it is possible that some of the editing sites are not translated or do not produce a stable protein, possibly due to deleterious effect of editing on the protein structure.

Protein Recoding Accounts for a Sizable Fraction of ADAR Editing in Neural Tissues of Octopus bimaculoides

For most organisms, A-to-I editing is markedly depleted from the protein-coding regions of the transcriptome. The question then arises of whether the extensive recoding in cephalopods is accompanied by extra-ordinary editing of the non-coding transcriptome. Recently, a genome was published for Octopous bimaculoides, the first from a cephalopod (Albertin et al., 2015), allowing us to use genome-dependent methods (Picardi and Pesole, 2013; Ramaswami et al., 2012) to study the full editome, including editing in non-coding sequences, as well as to compare with the genome-free method for the coding regions. Analyzing RNA-seq data of the same four neural tissues studied in the transcriptome-based approach resulted in 800,941 editing sites, 105,380 of them in annotated coding sequences (compared to 76,862 sites in coding sequences identified by the genome-free pipeline); 49,483 of these were also found using the transcriptome-based genome-free approach (Figure 3A). Differences between the two methods are due to the different de novo transcriptomes used and the different methods employed to filter

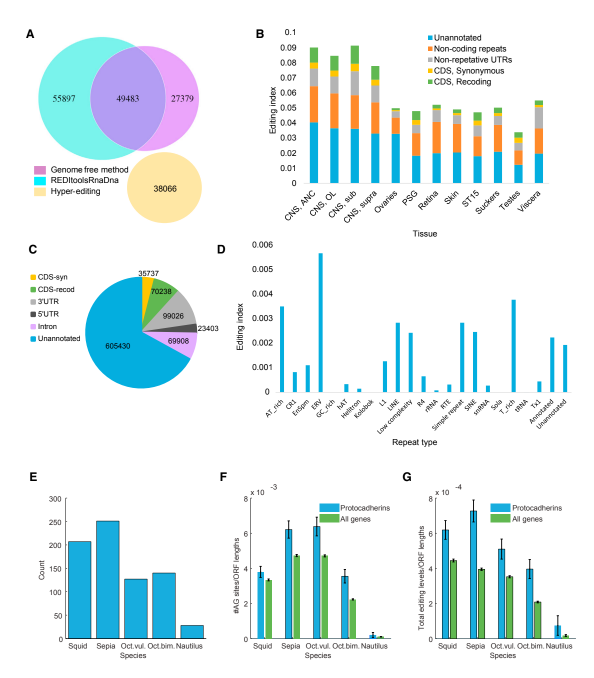


Figure 3. Editing in Octopus bimaculoides

(A) A-to-I editing sites were found within coding sequences of *Octopus bimaculoides* using three methods: the genome-free method (alignment to de novo transcriptome), the genome-dependent approach using REDItools (Picardi and Pesole, 2013), and identification of hyper-edited reads (Porath et al., 2014). Overall, the three methods identified 170,825 unique AG sites in *Octopus bimaculoides* coding sequences (38,066 hyper-editing sites do not overlap those found by the other methods). See STAR Methods for analysis of the differences between the results of the first two methods.

(B) RNA-editing levels, measured across the whole transcriptome (see Table S5) by the editing index (weighted average of editing levels over all editing sites identified in the transcriptome, see STAR Methods). Levels vary across tissues and are highest for neural tissues (see Table S6). Unlike mammals, a sizable fraction of editing events (11%–13% in neural tissues) result in recoding events. Annotation of transcripts and repeats is based on Albertin et al. (2015). (CNS = central nervous system; ANC = axial nerve cord; OL = optic lobe; Sub = subesophageal ganglia; Supra = supraesophageal ganglia; PSG = posterior salivary gland; ST15 = stage 15 embryo.)

- (C) The number of editing sites in coding region is comparable to the number found in introns.
- (D) Unlike the case in mammals, editing is not exceptionally enriched in specific repeat families in *Octopus bimaculoides*, as measured by the editing index (here defined as the editing level averaged over all, edited and unedited, adenosines in each specific repeat family).
- (E) Protocadherins is a gene family known to be principally expressed in the brain, important for mediating combinatorial complexity in neuronal connections and thought to play a role in diversifying neural circuitry (Chen and Maniatis, 2013). It was impressively expanded in Octopus bimaculoides (Albertin et al., 2015).

(legend continued on next page)

out random mismatches (see STAR Methods). These results suggest that the genome-free method provides a reasonable coverage of the editing signal in coding sequence, and that the number of editing sites outside the coding region is likely to be an order of magnitude higher than the number within the coding sequences for the other cephalopods studied here.

We analyzed RNA from 12 different tissues, including nonneural ones, and found 903,742 editing sites in the transcriptome (Table S5), 12% of which reside in coding regions (Figures 3B and 3C). In mammals, editing mostly occurs within genomic repeats (Bazak et al., 2014a; Levanon et al., 2004; Neeman et al., 2006). In primates specifically, most RNA-editing sites are found in Alu repeats, whose sequence facilitates the creation of a double-stranded RNA structure that promotes ADAR binding. Similarly, editing in Octopus bimaculoides is enriched in repeat regions (303,414/903,742 sites, 34%; 159,005 of them in annotated repeats). The "editing index," a robust measure of editing activity (Bazak et al., 2014b) defined as the editing level averaged over all adenosines (edited and unedited) weighted by expression level, is calculated to be 0.21% in octopus repeats for the panel of 12 tissues studied, which is comparable to the index observed in human Alu repeats (Bazak et al., 2014a). Unlike primates, though, there is not one specific repeat family that was found to contain the majority of sites, and SINEs are not edited more than other repeats (Figure 3D). Therefore, as the repeat editing index in octopus is calculated over all repeats (\sim 1.3 Gbp), and editing in repeats accounts for only 21%-38% of all editing events in octopus mRNAs (compared to >95% in primates), overall the number of editing events reflected in mRNA sequencing data is roughly an order of magnitude higher in Octopus bimaculoides compared to primates. Furthermore, in neural tissues $\sim 11\%$ –13% of these events result in amino-acid modification, compared with <1% in mammals (Bazak et al., 2014a). RNA editing is known to be important in neural function (Rosenthal and Seeburg, 2012), and abnormal editing patterns or ADAR function have been shown to underlie several neural conditions (Slotkin and Nishikura, 2013). Indeed, we find that editing in non-neural tissues of Octopous bimaculoides is roughly 2-fold lower, and recoding events are even more strongly suppressed (Figure 3B). Consistently, GO analysis of edited transcripts shows enrichment of neuronal and cytoskeleton functions in all four species (Table S6).

An intriguing result from the recently reported Octopus bimaculoides genome was that the protocadherin gene family was greatly expanded (Albertin et al., 2015). In the mammalian brain these proteins are important for mediating combinatorial complexity in neuronal connections and play a role in diversifying neural circuitry (Chen and Maniatis, 2013). We found a large number of protocadherins in the assembled transcriptomes for the four coleoid species (127-251 open reading frames [ORFs]), but not in nautilus (28 ORFs) (Figure 3E). Interestingly, protocadherins are significantly enriched in editing sites and are edited at higher levels in all four coleoid species, but not in nautilus (Figures 3F and 3G).

Signs for Positive Selection of Recoding Sites in Coleoid Cephalopods

Mammalian editing events in the coding region (and the editing levels) are negatively correlated with the importance of a site or gene-essential genes, and genes under strong functional constraints, tend to harbor lower numbers of editing sites and exhibit lower editing levels (Xu and Zhang, 2014). Furthermore, non-synonymous editing sites are suppressed, compared with synonymous ones, and the fraction of editing sites that are conserved across mammals is minute. These and other observations have led to the conclusion that although a few mammalian recoding sites are clearly beneficial, overall recoding by RNA editing is non-adaptive in mammals, presumably resulting from tolerable promiscuous targeting by the ADAR enzymes (Xu and Zhang, 2014).

To conduct a comparable analysis of the recoding repertoire in cephalopods, we first identified the non-synonymous editing sites. About 65% of cephalopod edits within coding sequences are non-synonymous, leading to 54,287-86,230 recoding sites in 6,688-8,537 ORFs (Table S7), orders of magnitude more recoding than any other species. In sharp contrast with mammals, thousands of recoding sites are shared between species (Figures 4A and 4B). As expected, the fraction of conserved sites is higher for species that are evolutionary closer (Figure 4C), but unlike the picture observed in other evolutionary lineages (Figure 4D), editing in coding sequences is, to a large extent, conserved. Interestingly, 1,146 editing sites (in 443 proteins) are conserved and shared by all four coleoid cephalopod species (Figure 4E). A large fraction of proteins are recoded at multiple sites, and many proteins harbor multiple conserved and highly edited (>10% in at least one species) recoding sites (Figure 4F; Table S8). Notably, even the editing levels in the shared sites are conserved and exhibit significant and sizable correlations between evolutionarily distant species (Figures 4G and S1).

Overall, the non-synonymous to synonymous (N/S) ratio for cephalopod edits is 65/35 = 1.9, as expected under neutrality taking into account the ADAR target motif (Alon et al., 2015). However, the N/S ratios increases to much higher values as editing levels increase (Figure 5A), signaling positive selection for the highly edited sites. Conserved sites show an even stronger pattern (Figure 5B), where almost all highly edited, conserved sites are non-synonymous. Consistently, and in stark contrast with mammals, the higher the editing levels, the more sites are conserved (Figures 5C and 5D). Furthermore, editing is over-represented in highly conserved regions of the transcriptome (>95% identity between species) (Figure 5E). Taken together, these results suggest that recoding by RNA editing is commonly adaptive in coleoid cephalopods, with many thousands of recoding sites under positive selection.

A large number of protocadherins are found in the assembled transcriptomes for the four coleoid species (127-251 open reading frames), but not in nautilus (28 open reading frames).

⁽F and G) Protocadherins contain significantly higher numbers of AG sites (F) and are edited at higher levels (editing level summed over all sites and normalized by ORF length), in all four coleoid species but not in nautilus (G). Error bars represent the SEM.

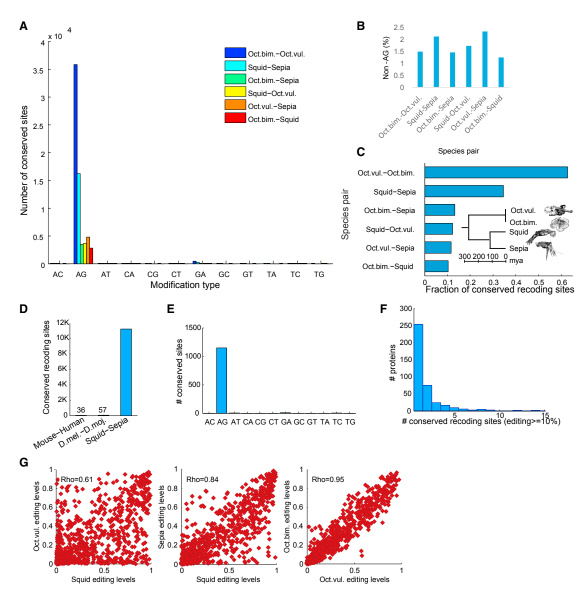


Figure 4. Extensive Recoding Is Conserved across Coleoid Cephalopods

(A) Tens of thousands of sites are conserved across species (see Table S7). The closer the species are evolutionarily, the higher the number of conserved sites. (B) Virtually all (97.5%–99%) mismatches conserved across species are A-to-G, resulting from A-to-I editing. Manual inspection of the few non-A-to-G mismatches appearing in multiple species suggests that they either result from systematic erroneous alignments or are actually editing sites that were mistakenly identified as G-to-A mismatches due to insufficient DNA coverage.

- (C) The majority of editing sites are conserved between the two octopus species, and even the most distant species share a sizable fraction of their sites.
 (D) In contrast, only 36 human recoding sites (1%–2% of human recoding sites) are shared by mouse, and a similar number are shared between *Drosophila melanogaster* and *D. mojavensis* (Yu et al., 2016) (diverged at later times than squid-sepia).
- (E) Interestingly, 1,146 AG modification sites (in 443 proteins) are conserved and shared by all four coleoid cephalopod species. Of these, 887 are recoding sites and 705 are highly edited (≥10% editing) recoding sites (in 393 proteins).
- (F) Some proteins include multiple highly edited recoding sites (see Table S8). Of note are uromodulin, α -spectrin (previously reported to harbor the highest number of recoding sites in squid; Alon et al., 2015), and calcium-dependent secretion activator 1 (CAPS1) with 14, 8, and 7 strong shared recoding sites, respectively. Recoding in CAPS1 was found to be conserved in vertebrate species from human to zebrafish (Li et al., 2009).
- (G) Not only are the locations of editing sites conserved, but their editing levels are correlated as well. Editing levels in 887 recoding sites shared by all species are highly, positively, and significantly correlated in all pairs of coleoid cephalopod species (p < 1e-75 for all pairs; see Figure S1 for three additional pairs). Correlation is higher the closer the species are to each other in evolutionary terms, with Pearson rho = 0.95 for the two octopus species.

Functional Implications of Recoding Sites

We next tested whether species-specific and conserved recoding events can affect protein function. We studied sepia, squid, and *Octopus vulgaris* K_v2 potassium channel orthologs, whose messages are abundantly edited (34–55 sites per species; 5 sites shared between all species; Figure S2 and Table S9).

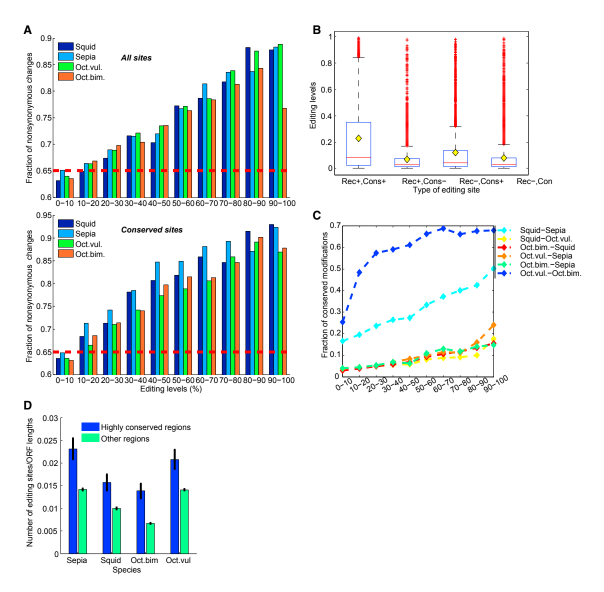


Figure 5. Signs for Positive Selection of Recoding by Editing

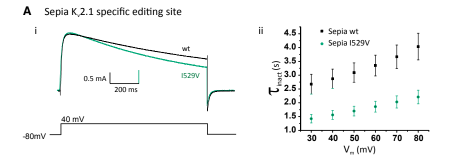
(A) The fraction of recoding sites among all editing sites in coding region increases with editing levels (top), as well as the fraction of recoding sites among all conserved sites (bottom). Red horizontal dashed line represents the recoding fraction expected assuming neutrality.

(B) Editing levels are higher in conserved recoding sites. Distributions of editing levels in four groups of putative A-to-I editing sites: recoding and conserved (Rec⁺, Cons⁺), recoding and non-conserved (Rec⁺, Cons⁻), conserved sites that cause a synonymous change (Rec⁻, Cons⁺), and non-conserved synonymous sites (Rec-, Cons-). Horizontal red lines mark the median level, and yellow diamonds mark the mean. Conservation and non-synonymity are both positively correlated with higher editing levels, as well as their interaction (ANOVA, p value < 1.0e-162). Data presented here for squid (conserved sites are conserved in sepia), but the results are similar and significant for all species.

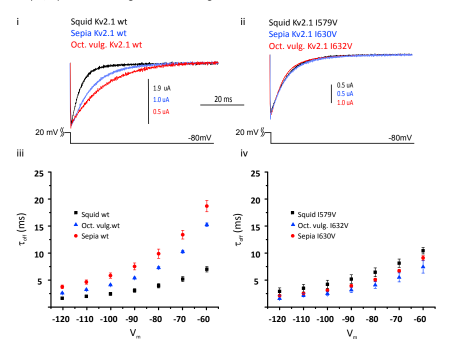
(C) In contrast with the case in humans, highly edited sites tend to be more conserved: the fraction of conserved sites rises with the editing level for all species pairs but more dramatically for the closely related octopuses and the sepia-squid pair.

(D) Highly conserved regions of the transcriptome are enriched in editing sites, further attesting to positive selection of RNA editing. Density of editing sites (number of AG sites normalized by length) is higher for 112 recoding regions that are highly conserved across the four species (> 95% identity; average length 1382 bp), compared with all other, less conserved regions (Wilcoxon p value < 0.001 for all species). Error bars represent the SEM.

Voltage-dependent potassium channels of the K_v2 subfamilly, also known as "delayed rectifiers," are expressed across the metazoa. In the mammalian central nervous system, they regulate excitability, action potential duration, and repetitive firing (Murakoshi and Trimmer, 1999). As with most voltage-dependent potassium channels, they are predominantly closed at negative membrane potentials and open at positive ones. When switched between negative and positive potentials, they open or close with characteristic rates. At positive potentials, channels will also spontaneously close after opening, a process known as "inactivation." The kinetics of these three processes play a vital role in determining how the channels regulate electrical signaling.



B Sepia, squid and oct. vulg. common editing site



To measure the effects of editing on functional properties, we expressed all channels in Xenopus oocytes and studied them using the Cut-Open Oocyte Vaseline Gap Voltage Clamp technique (Taglialatela et al., 1992). The unedited versions of the channels open over a similar range of voltages but have different opening, closing, and inactivation kinetics (Figures S3 and 6B). To examine the effects of editing, we first looked at the sepiaspecific editing site I529V. Figure 6Ai shows superimposed current traces, obtained in response to a voltage step from -80 mV to +40 mV, for the unedited and edited (I529V) versions of sepia K_v2. Clearly, the edited channel inactivates more quickly, at all voltages tested (Figure 6Aii; there is also a very modest slowing of channel closure upon bringing the voltage back to -80 mV). Editing had no effect on voltage sensitivity and channel opening (data not shown). We next looked at a common editing site (squid I579V, sepia I630V, and octopus I632V) and found that it predominantly affects the channels' closing rates. Interestingly, the direction of the effect is species dependent. First we analyzed the tail currents in the unedited squid, sepia, and octopus channels by recording currents at a negative membrane

Figure 6. Conserved and Species-Specific **Editing Sites Affect Protein Function**

Unedited (WT) and singly edited versions of the voltage-dependent K+ channels of the Kv2 subfamily were studied under voltage clamp (see Table S9).

(A) (i) Current traces resulting from a voltage step from -80 mV to 40 mV for the WT Sepia K₂2.1 and the same construct containing the sepia-specific I529V edit, lying within the 4th transmembrane domain (green), showing that I529V accelerates the rate of slow inactivation. (ii) Time constants for slow inactivation determined by fitting single exponentials to traces similar to those in panel (i) at different activating voltages (Vm).

(B) (i) Tail currents measured at a voltage (Vm) of -80 mV, following an activating pulse of +20 mV for 25 ms. Traces are shown for the WT K_v2.1 channels from squid, sepia, and Octopus vulgaris. (ii) Tail currents for the same channels edited at the shared I-to-V site in the 6th transmembrane span, following the same voltage protocol. (iii) Time constants from single exponential fits to tail currents obtained at various negative voltages (Vm) (following an activating pulse to 20 mV for 25 ms) show that the unedited channels close at distinct rates. (iv) but the edited versions close at similar rates. $N = 5 \pm SEM$ for all data plotted in this figure.

voltage of -80 mV following a brief activating pulse to a positive potential (Figure 6Bi). Each closes at distinct, species-specific rates, with sauid the fastest and octopus the slowest. However, upon introduction of the common editing event, the channels converge on a similar rate (Figure 6Bii); in squid, editing this site slows closing, whereas in octopus and sepia, it speeds it. This effect on closing kinetics was consistent at all voltages

tested (Figures 6Biii and 6Biv). Based on these data from K_v2 orthologs, and the fact that editing is exceptionally abundant in ion channels and proteins involved in synaptic vesicle release and recycling, the overall influence of RNA editing on neurophysiology is likely profound and complex.

Positive Selection of Editing Events Slows down Genome Evolution

To edit a specific adenosine within an RNA, ADAR enzymes require surrounding dsRNA structures. These structures are often large, spanning hundreds of nucleotides (Morse et al., 2002). If editing is under positive selection, maintaining these structures would require elevated sequence conservation in the vicinity of editing sites (Herb et al., 1996; Higuchi et al., 1993). As this sequence conservation stems from constraints related to RNA structure, rather than its coding capacity, it should affect synonymous and non-synonymous changes equally. Indeed, we see a marked depletion of inter-species mutations (Figures 7A and S4A-S4F) and intra-species genomic polymorphisms (Figures 7B and S4G), synonymous

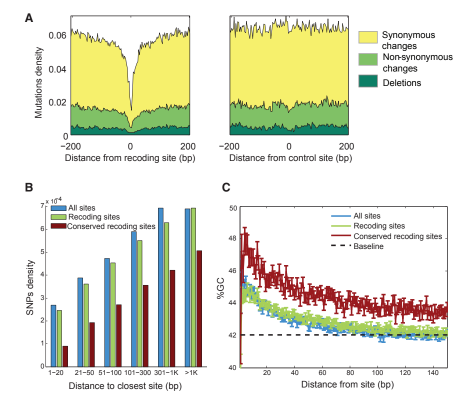


Figure 7. RNA Editing Slows down Cepha**lopod Genome Evolution**

(A) Inter-species mutations are purified from genome loci surrounding conserved recoding sites (data shown for sites shared by squid and sepia). Depletion of mutations extends up to \sim 100 bp of shared recoding sites (left). As a control, we show the mutation density (mutations/bp) around random non-edited adenosines from the same transcripts (right). Yellow-synonymous change; light green-non-synonymous; dark areen - deletions.

(B) Genomic polymorphisms are depleted near editing/recoding/conserved-recoding sites in squid, attesting to reduced genome plasticity. Effect is stronger for recoding sites, and even more so for the conserved recoding sites.

(C) GC content is elevated near editing sites in squid, allowing for more stable double-stranded RNA structures. The effect is even stronger in conserved sites. Dashed line represents the baseline GC level in the entire ORFome, and error bars represent the SEM.

See Figure S3 for analyses similar to those presented in (A)-(C) in other species.

and non-synonymous alike, up to ~100 nt on each side of a recoding site. These regions show an elevated GC content (Figures 7C and S4H), consistent with the requirement for the formation of stronger secondary structures.

The cumulative effect of this evolutionary constraint is considerable. Due to the large number of recoding sites and the extended range of the associated genomic rigidity, the local constraints observed in the vicinity of the recoding sites translate into a substantial global effect on genome evolution. These 200 nt windows around recoding sites cover a sizable fraction of all protein coding sequences: 23%-41%, depending on the coleoid species.

One may also quantify the effect of purifying selection in these regions by studying the fraction of inter-species mutations that were avoided, presumably due to maintaining the dsRNA structures required for editing. We analyzed the inter-species mutation rates (in orthologous parts of the respective transcriptomes) as a function of the distance to the closest conserved recoding site and found again that the rates are considerably lowered in the vicinity of editing sites, compared with the baseline rate observed far from any editing site (Figure S5A). Attributing the difference between the observed mutations rate and the baseline to effects of editing on genome evolution, and integrating this difference over the entire transcriptome, we estimate that 3%-15% of all transcriptomic inter-species mutations are purified (numbers vary for the various species pairs), apparently due to constraints imposed by editing. Similarly, we find that the actual number of SNPs in cephalopod coding sequences is 10%-26% lower than what would be seen in the absence of SNP suppression in the vicinity of recoding sites (Figure S5B). Thus, the purifying selection against inter-species mutations and intra-species genomic polymor-

phisms residing in proximity to recoding sites results in a sizable reduction in the global number of mutations and polymorphisms in these species, revealing an unanticipated genome rigidity required to maintain the extensive transcriptome recoding.

DISCUSSION

Seminal studies on RNA editing focused on recoding events and their functional outcomes (Burns et al., 1997; Higuchi et al., 1993). Later, with the advent of deep-sequencing technologies and accompanying computational advances, transcriptomewide screens showed that recoding is extremely rare. For example, there are millions of editing sites in the human transcriptome, but almost all of these reside in untranslated regions (Bazak et al., 2014a). This distribution implies some fundamental principles about RNA editing by ADARs. First, there is an active mechanism for excluding editing sites from coding regions; otherwise they would be far more common. Second, although there are clear exceptions for individual editing sites, the overall purpose of editing is not to recode (Liddicoat et al., 2015; Mannion et al., 2014). This point is reinforced by the fact that most mammalian recoding sites are neutral at best (Xu and Zhang, 2014). The abundant recoding in coleoids reported here runs contrary to these ideas.

We presented evidence that high-level recoding was invented by coleoids, or an extinct ancestor, after the divergence of the nautiloids. It is plausible that protein recoding may not be the primary function of editing in cephalopods. Perhaps there are other purposes for robust ADAR activity, such as its potential use in innate immunity (Liddicoat et al., 2015; Mannion et al., 2014).

As with any mutation, promiscuous "off-target" edits would sometimes be advantageous and therefore selected. However, many other organisms, such as humans, edit abundantly, producing multiple promiscuous edits. What is unique about coleoid cephalopods is that they appear not to exclude editing from protein-coding regions, leading to many thousands of recoding sites being recruited and conserved across distant species. Regardless of the primary motivation for editing, this unique phenomenon clearly has an enormous effect on the proteome.

The extensive recoding activity in cephalopods might suggest that there are underlying mechanistic novelties in their editing process, compared with other organisms. For example, cephalopod ADARs may have evolved to increase their catalytic activity or decrease their specificity. Previous studies have shown that squid express a splice variant of ADAR2 with an extra dsRNA-binding domain, and this feature increases the variant's affinity for dsRNA, leading to higher activity (Palavicini et al., 2009, 2012). Although cephalopods do express ADAR1 orthologs (Albertin et al., 2015; Alon et al., 2015), no functional studies have been conducted on them, nor on any invertebrate ADAR1 for that matter. They too may possess unique activities. Finally, one might expect the introduction of thousands of editing sites to be accompanied by undesirable side effects. For example, messages that contain so many mutations might often translate into dysfunctional, or even toxic, proteins. To accommodate this burden, cephalopods may have evolved unique mechanisms for protein folding and quality control. These ideas require further study.

Recoding in coleoid cephalopods is something of an enigma. Unlike the case for mammals, inter-species conservation and the higher-than-expected frequencies of non-synonymous changes suggest that a sizable fraction of events were recruited during the course of cephalopod evolution. Why would the coleoids choose to alter genetic information within RNA rather than hardwire the change in DNA? There are several potential advantages to making changes within RNA. First of all, the changes are transient. Thus an organism can choose to turn them on or off, providing phenotypic flexibility, a quality that is particularly useful for environmental acclimation (Garrett and Rosenthal, 2012; Rieder et al., 2015). In addition, RNA-level changes can better augment genetic diversity. With DNA, an organism is limited to two alleles. With RNA, all messages need not be edited, and thus the pool of mRNAs can include edited or unedited versions at given sites. When a message contains more than one site, complexity can increase exponentially. Future proteomic experiments will be necessary to determine whether the combinatorial complexity is realized in neural proteins, and whether editing contributes to neuron-specific diversity or the ability of the nervous system to respond to environmental cues. If the thousands of editing sites do indeed lead to independent functional outcomes, then the regulation of the editing process would be necessarily complex.

Among invertebrates, the nervous system of coleoids is uniquely large and complex. For example, with half a billion neurons, *Octopus vulgaris* has \sim 5 times the number of a mouse (Young, 1971). Coleoids have brain lobes dedicated to learning and memory (Hochner et al., 2003; Shomrat et al., 2008, 2015; Young, 1961) and exhibit a range of complex and plastic behav-

iors. Nautiloid brains are simpler, containing fewer neurons, and lack specific lobes dedicated to learning and memory (Young, 1965). The association of massive recoding with the nervous system, and the fact that it is unique to the coleoids and not observed in nautilus, hint at its relationship with the exceptional behavioral sophistication of the coleoids. This idea is reinforced by the high density of editing in transcripts that encode proteins directly involved in excitability.

What is most surprising about cephalopod recoding is its effect on genome evolution. From a mechanistic standpoint, this makes sense. In order to edit a specific adenosine, ADAR requires surrounding RNA structures. Even single nucleotide substitutions within these structures can abolish editing (Reenan, 2005; Rieder et al., 2013). If an editing site is advantageous, the structure must be preserved. Abundant editing requires abundant structures that can span a large fraction of the genomic coding sequence. Thus, although extensive recoding presents the species with a route toward proteome complexity, it comes with its own price tag. The constraints required to preserve thousands of recoding sites reduce the accumulation of mutations at positions in the proximity of an editing site, slowing down the rate of conventional, DNA-level evolution. The nervous system is one of the most important targets for natural selection, as subtle changes can lead to behavioral advances. For coleoid cephalopods, the need to make specific A-to-I changes within the neural transcriptome is sufficiently important to forego standard pathways of neuronal evolution.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

025#mmc7.

Supplemental Information includes five figures and ten tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2017.03.025.

An audio PaperClip is available at http://dx.doi.org/10.1016/j.cell.2017.03.

AUTHOR CONTRIBUTIONS

J.J.C.R. and E.E. conceived the study and designed the experiments, with the help of E.Y.L. and N.L.-B. N.L.-B. analyzed the data. S.A. developed some of the scripts used for analysis. H.T.P. did the hyper-editing analysis. J.J.C.R. performed all experiments, except for the MS, done by T.Z. and A.A. B.E. and R.U. analyzed the MS data. E.E. supervised the bioinformatics work, with input from E.Y.L. and J.J.C.R. N.L.-B., J.J.C.R., and E.E. wrote the paper.

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Biological Samples	0001102	DERTH LET
Octopus vulgaris	captured in San Juan, Puerto Rico	N/A
Sepia officinalis	Dr. Roger Hanlon, MBL, Woods Hole	N/A
Nautilus pompilius	SeaDwelling Creatures, Los Angeles	N/A
Critical Commercial Assays		
TruSeq DNA Sample Prep kit	Illumina	FC-121-3001
Truseq Stranded mRNA kit	Illumina	RS-122-2101
Deposited Data		
Squid DNA and RNA sequencing data	Alon et al., 2015	SRA, PRJNA255916
Octopus bimaculoides DNA and RNA sequencing data	Albertin et al., 2015	SRA, PRJNA270931, PRJNA285380
Sepia, Octopus vulgaris, Nautilus, DNA and RNA sequencing data	this paper	SRA, PRJNA299756
Aplysia DNA and RNA sequencing data		SRA, PRJNA13635, PRJNA77701
Mass spectrometry data	this paper	PRIDE repository, http://www.ebi.ac.uk/pride/database identifier: PXD005827
Experimental Models: Organisms/Strains		
Xenopus laevis (oocytes)	Xenopus Express, Inc, PO Box 10626. Brooksville, FL	IMP XL FM
Recombinant DNA		
Octopus vulgaris Kv2.1, see Table S10	IDT-DNA synthesized the channel; vector is from Liman et al., 1992	pOVKv2.1
Sepia oficianalis Kv2.1, see Table S10	IDT-DNA synthesized the channel; vector is from Liman et al., 1992	pSOKv2.1
Doryteuthis pealei Kv2.1, see Table S10	IDT-DNA synthesized the channel; vector is from Liman et al., 1992	pDPKv2.1
Software and Algorithms		
Trinity de novo transcriptome assembly	Grabherr et al., 2013	https://github.com/trinityrnaseq/trinityrnaseq/wiki Version: Trinity-r2012-10-05
Bowtie2 read alignment program	Langmead and Salzberg, 2012	http://bowtie-bio.sourceforge.net/bowtie2/index.shtml
REDItools editing detection package	Picardi and Pesole, 2013	https://sourceforge.net/projects/reditools/V-1.0.3
GOrilla functional analysis	Eden et al., 2009	http://cbl-gorilla.cs.technion.ac.il/
OrthoMCL - identification of orthologous ORFs	Li et al., 2003	http://orthomcl.org/orthomcl/
Proteome Discoverer	Thermo Fisher Scientific Inc.	http://www.thermofisher.comV 1.4
Other		
SwissProt database	Bairoch et al., 2005	http://www.uniprot.org/
De novo transcriptomes	this paper	http://www.tau.ac.il/~elieis/squid/

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Joshua Rosenthal (jrosenthal@mbl.edu), Eugene Bell Center, Marine Biological Laboratory, Woods Hole, MA 02543, USA.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

An adult male Octopus vulgaris was captured in October of 2014 at the mouth of the Laguna de Condado, San Juan, Puerto Rico. An adult male Sepia officinalis was provided by Dr. Roger Hanlon from the Marine Biological Laboratory in Woods Hole. This individual was raised in the Marine Biological Laboratory's Marine Resource Center from a fertilized egg that was collected in the English Channel off of Christchurch, Dorset England in 2014. The Stellate ganglia (SG), the optic lobe (OL) and a portion of the sperm sack were manually dissected from these specimens. One sample of Nautilus pompilius originating from the Philippines was obtained from "SeaDwelling Creatures" in Los Angeles. The optic lobe and the supraesophogeal ganglia were manually dissected from the sample. All samples destined for RNA extraction were immersed in chilled, filtered seawater and immediately preserved in RNA later. The samples intended for DNA extraction were flash frozen in liquid nitrogen. All samples were then stored at -80° C. RNA from all tissues was extracted with the RNAqueous kit (Life Technologies, Carlsbad, CA), and genomic DNA was extracted from the sperm sack using Genomic Tip Columns (QIAGEN, Venlo, Limburg, the Netherlands).

We added previously described RNA and DNA sequencing data from the squid species Doryteuthis pealeii (Alon et al., 2015) (PRJNA255916), as well as published data from another Octopus species, Octopus bimaculoides (PRJNA270931, PRJNA285380), whose genome was recently sequenced (Albertin et al., 2015). We also added available RNA and DNA samples from the mollusk Aplysia californica, as an evolutionary outgroup (PRJNA13635, PRJNA77701).

METHOD DETAILS

Library preparation and sequencing

The genomic DNA sequencing library for Octopus vulgaris, sepia and nautilus were prepared using the TruSeq DNA Sample Prep kit, as described by the manufacturer (Illumina, San Diego, CA), and sequenced using three lanes of the Illumina HiSeq 2000 instrument. The RNA-Seq libraries for all the samples were prepared using the TruSeq Stranded mRNA Sample Prep Kit, as described by the manufacturer (Illumina), and were sequenced using one lane for each sample of Illumina HiSeq 2000 instrument.

llumina sequencing was utilized to generate paired-end, 151 nt reads, using RNA from OL and SG tissues for Octopus vulgaris and sepia, and OL and subesophageal ganglia for nautilus. For DNA sequencing, 101 nt reads were produced. The number of reads generated for each tissue is presented in Table S1.

Editing site validation

Validation of editing sites predicted by our bioinformatics pipeline was previously performed and reported (Alon et al., 2015). In brief, direct Sanger sequencing confirmed editing at 40/40 A-to-G squid recoding sites, and deep-sequencing validated 120/143 A-to-G recoding sites but none of the 12 non A-to-G sites tested. In this work we take validation a step further and examine editing at the protein level.

In-gel proteolysis and mass spectrometry analysis

We applied proteomic mass spectroscopy (MS) analysis to examine the extent to which these RNA modifications are translated into the proteome. Squid giant axon and stellate ganglion samples were separated by a SDS-PAGE and the gel was stained with Coomassie Blue and sliced to 5 slices. The slices were processed for tryptic digestion by first reducing the disulfides with 3 mM DTT in 100 mM ammonium bicarbonate for 20 min at 60°C. Next, the sulfhydryl were carboxymethylated with 10 mM iodoacetamide in 100 mM ammonium bicarbonate in the dark for 30 min at room temperature. The proteins were in-gel digested with modified trypsin (Promega) in 10% acetonitrile and 10 mM ammonium bicarbonate at a 1:10 enzyme-to-substrate ratio overnight at 37°C. Additional trypsinization was done for 4 hr.

The resulting tryptic peptides were desalted on C18, Stage-Tip (Ishihama et al., 2006) and resolved by reverse-phase chromatography on 0.075 X 200-mm fused silica capillaries (J&W) packed with reversed phase Reprosil-C18-Aqua (Dr. Maisch GmbH, Ammerbuch-Entringen, Germany) as in (Ishihama et al., 2002). The peptides were eluted with linear 105 min gradient of 5% to 28% acetonitrile with 0.1% formic acid in water, followed by 15 min gradient from 28% to 90% acetonitrile and 15 min at 90% acetonitrile at flow rates of 0.15 μl/minutes. MS was performed by a Q-Exactive-Plus mass spectrometer (Thermo Fisher Scientific) in a positive ion mode using repetitively full MS scan followed by Higher-energy Collision Dissociation (HCD) of the 10 most dominant ions, selected from the first MS scan. A dynamic exclusion list was enabled with an exclusion duration of 20 s.

The MS data was analyzed using Proteome Discoverer 1.4 software with the Sequest (Thermo Fisher Scientific) algorithm against the specific databases, combining all 5 fractions of each sample in one search. Minimal peptide length was set to six amino acids and a maximum of two miscleavages was allowed. Mass tolerance of 15 ppm for the precursor masses and for the fragment ions. Peptide- and protein-level false discovery rates (FDRs) were filtered to 1% using the target-decoy strategy. Semi quantitation was done by calculating the peak area of each peptide based its extracted ion currents (XICs), and the area of the protein is the average of the three most intense peptides from each protein.

Expression and recording of cephalopod K_v2 channels in *Xenopus* oocytes

Full-length Octopus vulgaris, Sepia and Squid K_v2.1 constructs, to be expressed in Xenopus oocytes, were based on the unedited amino acid sequences deduced from the transcriptome assemblies. The sequence for the two Octopus species are almost identical (Table S4), so we studied only one of them. Codon optimized versions for Xenopus laevis were synthesized using gene blocks and cloned into the Xenopus expression vector pGEMHE (Liman et al., 1992) using a Gibson Assembly. Single RNA-editing sites were added to these clones by standard oligonucleotide-based site-directed mutagenesis using the Quickchange Lightening Site-Directed Mutagenesis Kit (Agilent Technologies). Capped, polyA-tailed cRNA from each clone was transcribed using the T7 mMessage mMachine Kit (Thermo Fisher) and injected into stage V and V1 Xenopus oocytes at approximately 750 pg/oocyte. Currents were recorded 2-3 days after injections using the Cut-Open Oocyte Vaseline Gap technique (Lockery and Goodman, 1998), using a relatively slow sampling rate. The external solution contained: 20 mM K-Glutamate, 100mM N-Methyl-D-Glucamine-Glutamate, 2.5 mM MgCl2, 2.5mM CaCl2, 10 mM HEPES, pH = 7.4. Oocytes were permeabilized using 0.3% Saponin in the internal solution. Voltage was controlled using a CA-1B High Performance Oocyte Clamp (Dagan Corporation). Analog currents were digitized at 100 kHz, and voltage commands were made, using an SBC6711 A/D D/A board (Innovative Integration, Simi Valley CA). Signals were filtered at 5 kHz. To avoid series resistance errors, only traces exhibiting less than 10 mA current were used for analysis. Data collection and clamp command was made using GPATCH M software, kindly provided by Dr. F Bezanilla (University of Chicago). Leak currents were subtracted using a linear P/4 procedure. Data were analyzed using the ANALYSIS software, also provided by Dr. F. Bezanilla. Deactivation kinetics were measured by fitting a single exponential of the form $y = Ae^{-\tau/T} + B$ to the traces where A = current amplitude, τ = the time constant, T = time, and B = a constant baseline. Inactivation kinetics were measured by fitting a single exponential of the form $y = Ae^{-\tau/T}$ to the traces where A = current amplitude, $\tau = the$ time constant, and T = time. Relative conductance was measured using peak amplitude of tail currents. N = 6 for all data points ± SEM.

QUANTIFICATION AND STATISTICAL ANALYSIS

Transcriptome assembly

The species studied, with the exception of Octopus bimaculoides, do not have a reference genome. We therefore utilized the RNA-Seg data to assemble a transcriptome using the Trinity de novo assembly package, Version: Trinity-r2012-10-05 (Grabherr et al., 2013). We found that Trinity does not perform well for reads longer than 100bp, and therefore all reads were trimmed (symmetrically, on both sides) to this length prior to assembly. For consistency, we assembled the transcriptome for Octopus bimaculoides in the same manner, using data from the four neural tissues out of the 12 tissues available: optic lobe (OL), supraesophogeal ganglia (supra), subesophogeal ganglia (sub), and the axial nerve cord (ANC). For each gene, we kept only the longest isoform for downstream analysis. The assembly statistics are summarized in Table S2.

Detection of editing sites

In order to detect editing sites, using the matched RNA and genomic DNA samples, we employed a method similar to that described in (Alon et al., 2015). Briefly, RNA and DNA reads were separately aligned against the assembled transcriptome using Bowtie2 with local alignment configuration and default parameters (Langmead and Salzberg, 2012). Reads that were not uniquely aligned were discarded. We identified potential open reading frames (ORFs) in the assemblies by locating components that were found to be significantly similar (Blastx E-value < 1e-6) to the Swiss-Prot proteins dataset (Bairoch et al., 2005). Each ORF was extended until either a stop codon or the end of the Trinity component was met. Table S2 summarizes the properties of the ORFs found for each species. To detect editing events, we applied a binomial test to locate significant modifications between RNA reads and the Swiss-Prot ORFs and distinguish them from sequencing errors or SNPs. For more detailed description of the editing detection method, see (Alon et al., 2015). Two important modifications were introduced here with respect to the scheme presented in (Alon et al., 2015): (1) we discarded all mismatches that occur up to 6bp from alignments' ends (2) we discarded all reads that contained more than one mismatch type (e.g., A-to-G mismatch and A-to-C mismatch is the same read), or more than two mismatches (of any type) altogether, as these are suspected to be misaligned. The number of reads discarded ranges between 0.6 and 1.7 million, for the species studied. Of these, 200-800 thousand reads might have exhibited two editing events. Unlike (Alon et al., 2015), we did not distinguish between "weak" and "strong" sites. Rather, the genomic strand was determined by the DNA reads (and in the absence of DNA coverage, by the majority of RNA reads). In case of conflicting DNA reads, sites were discarded.

Editing in Octopus bimaculoides

Our genome-free detection scheme is limited by the quality of the assembled de-novo transcriptome, and the focus on coding sequences only. The recent sequencing of Octopus bimaculoides genome (Albertin et al., 2015) enabled us to compare our genome-free scheme to a genome-based one, obtain an independent assessment of the true extent of editing in coding sequences, and estimate the full picture of the octopus editome outside of coding sequences.

The REDITools package (V-1.0.3) (Picardi and Pesole, 2013) was used to locate RNA-DNA differences in the octopus genome, using RNA-seg data from the same four neural tissues that were used in the genome-free scheme. Editing sites annotation was based on the transcriptome provided in (Albertin et al., 2015), which differed from the one we used in the genome-free method.

We found 800941 AG sites, 105380 of them in annotated coding sequences (compared to 76862 sites in coding sequence identified by our transcriptome-based pipeline), 49483 of these sites were also found using our pipeline (see Table S5). Therefore, we see that our pipeline provides a reasonable coverage of the editing signal in coding sequence, and that the number of editing sites outside the coding region is likely to be an order of magnitude higher than the number within the coding sequence for the other cephalopods studied here. The differences between the two methods stem from the different de-novo transcriptome used, and the different parameters used to assess the mismatches observed (REDItools uses cutoffs on the number of reads, while the genome-free approach applied a binomial analysis). Using BLAST to compare the two transcriptomes, we find that \sim 90% of Octopus bimaculoides transcripts detected in our pipeline are covered by the transcriptome described in (Albertin et al., 2015), but only ~68% of the latter transcriptome described in (Albertin et al., 2015), but only ~68% of the latter transcriptome described in (Albertin et al., 2015). scripts are covered by our detected ORFs, which explains the majority of the sites missed by our pipeline. Of the \sim 56K sites found by REDItools but not by the genome-free method, 35911 (64%) are located within transcripts missing from our de-novo transcriptome and the rest are seen by the genome-free method but fail to achieve statistical significance (e.g., cases where coverage is very high, and only a few G's are observed). Out of the ~27K sites found using the genome-free method but not by REDItools, about 14k (52%) were missed due to too strict parameters employed by REDItools, and \sim 13K sites (48%) reside in sequences that do not exist in the transcriptome assembled in (Albertin et al., 2015).

In order to profile the editome in all available tissues, we re-did the analysis using all available RNA data from 12 tissues available: axial nerve cord (ANC), optic lobe (OL), subesophageal brain (Sub), supraesophageal brain (Supra), Ovaries, posterior salivary gland (PSG), retina, skin, stage 15 embryo (ST15), suckers, testes and viscera (heart, kidney and hepatopancreas), leading to a total of 903.742 sites.

As editing is so abundant in these organisms, it is expected that many sites will reside in reads that are extensively edited, to the point they are not aligned to the reference genome/transcriptome by the standard alignment tools. Thus, we also applied the method suggested in (Porath et al., 2014) to identify hyper-edited reads, leading to 38066 additional sites in coding regions, not found by the other methods (Figure 3A).

The editing index was used to compare editing activity across different tissues (Figure 3B). It is defined as the number of 'G's in RNA-reads nucleotides that were aligned to the predicted editing sites, divided by the total number of read-nucleotides that align to these positions ("A"s and "G"s). In order to compare editing between different repeats, we used the repeats editing index, calculated in the same way over all genomic adenosines within repeats (number of A-to-G mismatches in RNA-reads nucleotides that were found in repeats, divided by the total number of read-nucleotides that align to genomic adenosines in repeats). The higher the index, the more editing occurs in the specific repeat family element (Figure 3D).

Functional analysis of edited ORFs

To test for functional enrichment, we ranked the genes by cumulative editing levels (editing levels summed over all sites within the gene, normalized by ORF length), and used the online tool GOrilla (Eden et al., 2009). As a control, the genes were also ranked by expression levels measured by FPKM and analyzed in the same manner (Table S6).

Finding orthologous editing sites

We used OrthoMCL (Li et al., 2003) to identify orthologous pairs of ORFs, using only best-two-way hits. This covered 77% of the transcriptomes. Then, for each editing location in the query species of each pair, we screened for the best possible aligned region in the ortholog using BLAST alignment scores, and found the matching nucleotide and amino acid. A conserved editing site is a case where the same mismatch occurs at the exact orthologous nucleotide, leading to the same amino acid substitution.

Identifying SNPs

In order to minimize false positives, we took a conservative approach, and called a genomic location a SNP if the DNA reads aligned to this location show exactly two types of nucleotides, with at least five reads supporting each type, and each of the types being supported by 30%-70% of the reads. Obviously, this scheme is not meant to exhaust the list of SNPs in these species, which is anyway impossible using a single animal per species, but suffices to estimate the relative depletion of SNPs around recoding sites.

DATA AND SOFTWARE AVAILABILITY

The data used in the study are publicly available at the Sequence Read Archive (SRA), accessions PRJNA300723. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE (Vizcaíno et al., 2016) partner repository with the dataset identifier PXD005827. The de-novo constructed transcriptomes used for the analysis are available at http://www.tau.ac.il/~elieis/squid.

Supplemental Figures

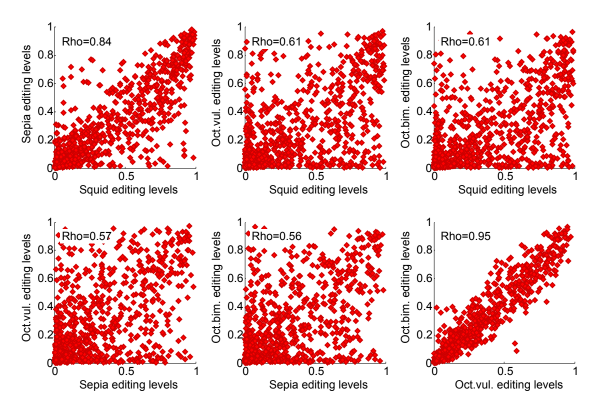
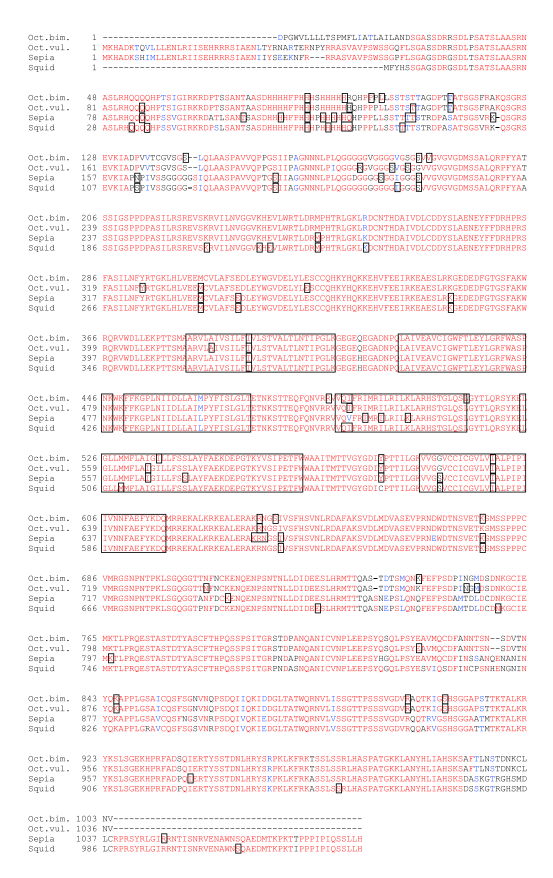


Figure S1. Editing Levels of Shared Sites Are Correlated across Coleoid Species, Related to Figure 4

Editing levels in 887 recoding sites shared by all species are highly, positively and significantly correlated in all pairs of coleoid cephalopod species (p < 1.0e-75 for all pairs). Correlation is higher the closer the species are to each other in evolutionary terms, with Pearson rho = 0.95 for the two octopus species.





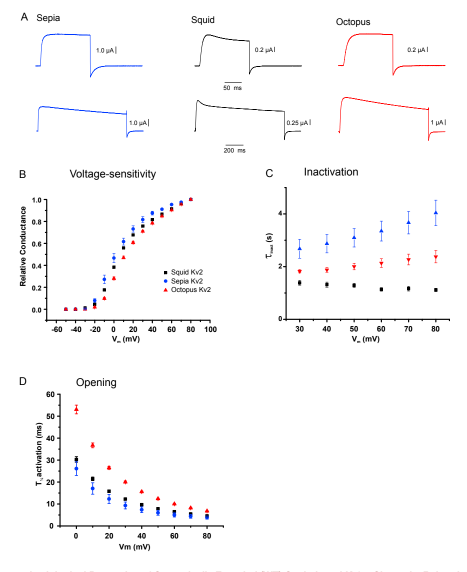


Figure S3. Basic Electrophysiological Properties of Genomically Encoded (WT) Cephalopod $K_{\nu}2.1$ Channels, Related to Figure 6

- (A) Examples of current traces recorded on fast (top) and slow (bottom) time-bases following activating pulses to +60 mV from a holding potential of -80 mV. These traces illustrate the variable activation and inactivation rates of the genomically encoded channels.
- (B) The channels' voltage sensitivities are similar. Relative conductance was measured from normalized tail-current amplitudes following 150 ms activating pulses to various activation voltages (V_m) .
- (C) Inactivation time constants (τ) taken from single exponential fits to current traces measured during a 1 s activation step of V_m (similar to the traces presented in panel A).
- (D) The half-times for 150 ms activating pulses were measured from traces similar to those presented in panel A. They measure the time it takes for the currents to reach one half of their maximum values following an activating voltage step. $N = 6 \pm SEM$ for all graphs.

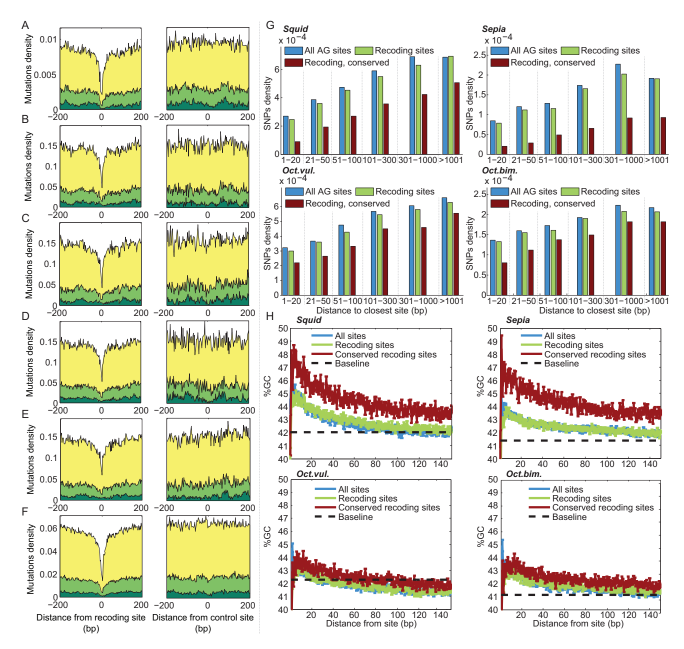


Figure S4. Reduction in Intra- and Inter-Genomic Variability around Editing Locations, Related to Figure 7

(A–F) Species-to-species mutations are suppressed within ~100 bp of a recoding site shared by the two species (left panels). No effect is seen when taking random non-edited sites from conserved regions within the same transcripts (right panels) (A) Oct.bim.-Oct.vul. (B) Oct.bim.-Sepia (C) Oct.bim.-Squid (D) Sepia-Oct.vul. (E) Squid-Oct.vul. (F) Squid-Sepia. (yellow – synonymous change, light green – non-synonymous; dark green – deletions; Mutations density is the number of mutations found at a given distance from an editing site divided by the number of such sites, i.e., mutations per base-pair).

(G) Intra-species genomic polymorphisms are suppressed in genomic loci surrounding editing sites. Genomic polymorphisms are depleted near editing/recoding/conserved-recoding sites, attesting for a decrease in genome plasticity. Effect is stronger for recoding sites, and even more so for the conserved recoding sites.

(H) Higher GC content in the vicinity of editing sites. GC content is elevated near editing sites. This effect persists up to \sim 100 bp, and is therefore unlikely to be explained by the local ADAR preferences. Rather, it may facilitate formation of stronger and more stable double-stranded RNA structures required for ADAR binding and editing. Blue: all editing sites; green: all recoding sites; red: conserved recoding sites; dashed black line: GC content averaged over all ORFs. Error bars represent the SEM. The effect is not seen in for AG mismatches found in nautilus, expected to be mostly false-positives.

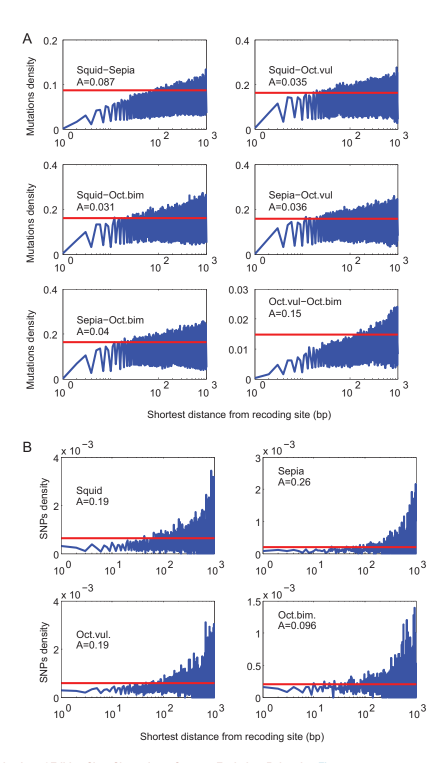


Figure S5. Positive Selection of Editing Sites Slows down Genome Evolution, Related to Figure 7

(A) Mutation rates (averaged over all orthologous parts of the transcriptome) are presented as a function of the distance of the genomic base to the closest conserved recoding site. The baseline mutation rate (red) represents the rate measured for all sites at distances > 1,000 bp from a recoding site (including mutations in ORFs harboring no editing site). Integrating the difference between the baseline and the observed rate for genomic location at a distance shorter than 1,000 bp provides an estimate of the number of avoided mutations, apparently due to the constraints imposed by editing. We estimate the fraction of avoided mutations as the number of avoided mutations divided by (#avoided+#observed), and find this fraction (marked as A in the different panels) to be 3%–15%. This is an underestimation, as sites not conserved between the two species, or sites in transcriptome regions not conserved between the two species, are not taken into account.

(B) Similar to the analysis presented in (A), we looked at the SNPs density (number of SNPs per bp) as a function of the distance of the genomic base to the closest recoding site. The baseline SNPs density (red) is the density observed at distances of > 500 bp from a recoding site (including SNPs in ORFs harboring no editing site). Integrating the difference between the baseline and the observed rate for genomic location at a distance shorter than 500bp provides an estimate of the number of avoided SNPs, apparently due to the constraints imposed by editing. Here too, we estimate the fraction of avoided SNPs as the number of avoided SNPs divided by (#avoided+#observed), and find this fraction (marked as A in the different panels) to be 10%–26%. This is an underestimation, as the contributions of sites in non-conserved transcriptome regions, are not taken into account.