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Survey of 800+ data sets from human tissue and body fluid reveals xenomiRs are likely artifacts

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ABSTRACT
miRNAs are small 22-nucleotide RNAs that can post-transcriptionally regulate gene expression. It has been proposed that dietary plant miRNAs can enter the human bloodstream and regulate host transcripts; however, these findings have been widely disputed. We here conduct the first comprehensive meta-study in the field, surveying the presence and abundances of cross-species miRNAs (xenomiRs) in 824 sequencing data sets from various human tissues and body fluids. We find that xenomiRs are commonly present in tissues (17%) and body fluids (69%); however, the abundances are low, comprising 0.001% of host human miRNA counts. Further, we do not detect a significant enrichment of xenomiRs in sequencing data originating from tissues and body fluids that are exposed to dietary intake (such as liver). Likewise, there is no significant depletion of xenomiRs in tissues and body fluids that are relatively separated from the main bloodstream (such as brain and cerebro-spinal fluids). Interestingly, the majority (81%) of body fluid xenomiRs stem from rodents, which are a rare human dietary contribution but common laboratory animals. Body fluid samples from the same studies tend to group together when clustered by xenomiR compositions, suggesting technical batch effects. Last, we performed carefully designed and controlled animal feeding studies, in which we detected no transfer of plant miRNAs into rat blood, or bovine milk sequences into piglet blood. In summary, our comprehensive computational and experimental results indicate that xenomiRs originate from technical artifacts rather than dietary intake.

Keywords: miRNA; microRNA; meta-study; exogenous miRNAs; next-generation sequencing; contamination; artifacts

INTRODUCTION
Over the past decade, RNAs extracted from cell-free body fluids have received increasing attention as novel diagnostic markers for disorders as diverse as narcolepsy, cancers, and viral infections (Mitchell et al. 2008; Winther et al. 2013; Holm et al. 2014a). Many distinct body fluids have been profiled for biomarkers (Weber et al. 2010; Holm et al. 2014b). Blood samples are most commonly used, as they are obtained from the human body with minimal invasion and can give snapshots of the transcriptome of many tissues that are communicating with the bloodstream (Chen et al. 2008; Schwarzenbach et al. 2014). Profiling of microRNAs (miRNAs), a group of small noncoding RNAs, is particularly promising because the only ∼22 nucleotides (nt) short fully matured miRNAs are considerably stable compared to other RNA species, such as mRNAs (Jung et al. 2010). The maturation is a process of sequential endonucleolytic cleavages of the primary miRNA transcript by RNase III enzymes (Ha and Kim 2014). After cleavage the mature miRNA is incorporated into the miRISC complex, where it serves to guide the complex to target sites in mRNAs. Once bound, the complex can regulate mRNA gene expression...
by inhibiting translation, promoting degradation or relocation to subcellular foci where mRNA is no longer accessible to the translational machinery. miRNAs in human body fluids have been identified as emerging effective biomarkers in many different biological states as well as diseases (Simonson and Das 2015; Fabris and Calin 2016). While most cell-free body fluid biomarker studies have been conducted by conventional quantitative PCR-based methods, an increasing number of more recent studies use deep sequencing as a method that is independent of prior knowledge of transcript sequence composition (Freedman et al. 2016).

The current state-of-the-art method for profiling body fluid miRNAs is next-generation sequencing of small RNAs (sRNA-seq). This technique detects millions of miRNAs in a single experiment and allows discovery of new transcripts (Burgos et al. 2014; Byron et al. 2016). Sequencing allows detection of miRNAs in an unbiased way that can lead to surprising results. For example, a recent report from Zhang et al. (2012) suggested that a miRNA (osa-MIR168) from a rice-rich diet can cross the human gut epithelium and enter the bloodstream where it can interact with and repress the human high-density lipoprotein RAP1 transcript. The ramifications of this observation are profound, since it implies that cross-species miRNAs (“xenomiRs”) can impact the human transcriptome through diet. This could open up a new research field of nutrition-specific RNA-based remedies and development of oral delivery of RNA-based medicine.

The xenomiR hypothesis is indirectly supported by studies showing that miRNAs extracted from plants (Philip et al. 2015) or from body fluids (Chen et al. 2008) are stable when subjected to high temperature or enzymatic digestions. Some in vivo studies support the initial finding that xenomiRs are present in body fluids of humans and other animals, however at a very low expression level (Wang et al. 2012, 2013). The most highly abundant Brassica oleracea MIR172 was found in stomach, intestine, serum, and feces of mice that were fed plant RNA extracts. The MIR172 abundance and peak time in multiple origins of samples was consistent with ordered dietary update: stomach > intestine > feces > serum (Liang et al. 2014). Another study found several plant and insect xenomiRs in serum and plasma samples. Interestingly, the rice-derived xenomiRs were dominating in the serum sample from Chinese ethnicity compared to the plasma sample from Caucasian ethnicity, which was dominated by corn-derived xenomiRs (Wang et al. 2012). In a recent study, new evidence of xenomiRs was presented showing that food-derived miRNAs were transferred to mammalian placenta and directly regulating fetus gene expression (Li et al. 2015). In another mouse study, the authors reported that MIR2911, derived from the honeysuckle plant, was present in animals drinking or gavage feeding honeysuckle. Additionally, the authors showed that the miRNAs were able to regulate influenza virus by reducing replication and rescued viral infection (Zhou et al. 2015).

In contrast, several studies were unable to reproduce the original findings or have provided direct evidence that the xenomiRs are the result of technical contaminations rather than diet-dependent cellular uptake. A number of groups have searched for plant xenomiRs in human body fluids but have been unable to find these sequences at any substantial level (Snow et al. 2013; Witwer et al. 2013). Similarly, other groups were unable to detect dietary xenomiRs when conducting controlled feeding studies in humans, mice, bees, and other insects (Dickinson et al. 2013; Witwer et al. 2013; Auerbach et al. 2016). Further, one independent research group investigated the xenomiR composition of sRNA-seq data from amphioxus animals, generated by the team of the original Zhang et al. (2012) publication. It was found that the amphioxus data contained rice miRNAs in exactly the same composition as in the previous human serum study from the same group (Tosar et al. 2014). Given that amphibia have an exclusive algae-based diet, this strongly suggests that the rice miRNAs identified in both studies are the result of contamination. In general, contaminations can occur at several steps in sequence data generation. They can happen if traces of samples are mixed, either in the field, clinic, or laboratory. Since next-generation sequencing is tremendously sensitive, a few molecules could feasibly cause false detection of xenomiRs. Functional contaminations can also occur during sequencing or the data analysis. Often, several sample cDNA libraries are sequenced together, and each read is assigned back to the sample through a barcode. If cDNA molecules are in spatial proximity during sequencing, their respective barcodes can be swapped. Additionally, if the barcodes are similar in sequence, the defaults analysis software can misassign the reads to the wrong sample (Kircher et al. 2012).

Despite the many studies on xenomiRs, it still remains unresolved if they originate from diet or from laboratory contaminations. Public sequence repositories contain hundreds of sRNA-seq data sets from human tissues and body fluids. Surprisingly, this resource has not been mined in a systematic manner to study the presence and compositions of xenomiRs. To address this, we conducted the first comprehensive meta-study of xenomiRs in 824 public human data sets. We developed a new computational pipeline to standardize and normalize the raw sequence data, allowing us to compare the results from multiple studies. We found that xenomiRs are present in many human sRNA-seq data sets, although in very low abundances. There is no enrichment of xenomiRs in samples that are exposed to dietary intake (liver and serum) compared to samples that are protected by the blood–brain barrier (brain and cerebrospinal fluid, CSF). The body fluid samples are dominated by xenomiRs from rodents, which are rare in human diets, but are common laboratory animals. We observe that the body fluid xenomiR profiles group by study (batch effects), suggesting that technical factors influence the xenomiR presence. In complementary controlled animal feeding studies, we see no transfer of dietary plant miRNAs into the bloodstream of rats, nor of cow milk miRNAs into piglets. In summary, our computational and wet-lab results provide evidence for
Meta-study reveals xenomiRs are likely artifacts

RESULTS

Meta-study of 824 human sRNA-seq data sets

We collected 824 public human sRNA-seq data sets, comprising a total of 10 billion sequencing reads, to profile the presence and abundance of xenomiRs (Supplemental Table 1). These data sets included 215 serum and plasma samples and 37 exosome samples, since these are body fluids that have previously been described to contain xenomiRs (Wang et al. 2012, 2013; Zhang et al. 2012; Liang et al. 2015). We also selected 180 samples of cerebrospinal fluid (CSF), which is separated from the bloodstream by the brain–blood barrier (BBB), which unprotected small RNAs are unlikely to cross (Pardridge 2005; Boudreau et al. 2011). To investigate whether xenomiRs accumulate in cellular tissues, we selected 93 samples from liver and 197 samples from blood cells. We considered these to be the tissues most likely to be exposed to dietary transcripts. As a negative control, we selected 102 samples from human brain, since this organ is separated from the bloodstream by the BBB. The raw sequence data from the 824 sets were selected, quality controlled, and preprocessed using a standardized computational pipeline, to ensure that results are comparable. Specifically, the data sets were selected and automatically downloaded from the public GEO database based on keyword searches (Fig. 1; Supplemental Fig. 1; Materials and Methods). The data were subjected to manual inspection and a subsequent stringent computational quality control based on sequence quality and complexity. The remaining reads were preprocessed by removing the sequencing adapters and discarding short reads that are unlikely to be functional miRNAs. The processed data were then used to assess the presence and abundance of xenomiRs.

XenomiRs are largely absent in brain, liver, and blood cell samples

We first investigated the presence of xenomiRs in the 392 human tissue data sets, comprised of samples from liver, brain, and blood (we here consider blood cells and whole blood as tissue). To perform a stringent analysis, we only considered miRNAs that are specific to certain animal and plant clades (branches of the phylogenetic tree) (Sempere et al. 2006; Wheeler et al. 2009; Taylor et al. 2014). For instance, the miR-3 family is known to be present only in insects, so the presence of miR-3 constitutes strong evidence for insect RNA. In contrast, deeply conserved miRNAs like let-7 are present in all bilaterian animals. Therefore, it is difficult to distinguish insect let-7 and primate let-7, given that sequence differences can be obscured by RNA editing, sequencing errors, etc. A full list of the conservative clade-specific miRNA families can be found in Supplemental Table 2. When analyzing the human 392 tissue samples, we observed abundant primate-specific miRNAs in all samples (Fig. 2). This is according to expectations, as these very likely represent bona fide endogenous human miRNAs. On average, a median of 979 primate miRNA counts from 17 distinct primate miRNA families were detected (Supplemental Fig. 2A,D). It should be noted that these counts are substantially less than the total number of human miRNAs, since we here only consider the primate clade-specific miRNAs, which tend to be poorly expressed. While primate-specific miRNAs were found in 100% of the samples, xenomiRs (combined clade-specific miRNAs from rodents, insects, etc.) were present in only 17% of the tissue samples (Supplemental Fig. 3A). Some entire studies were completely void of any xenomiRs, such as GSE67491 with 30 samples comprising more than 244 million reads. This complete absence of xenomiRs in some large studies demonstrates the high specificity and low false-positive rate of our pipeline. In the samples that did contain xenomiRs, we found a median of 3.5 xenomiR read counts and 1.5 distinct xenomiR families, much lower than for the primate-specific miRNAs (Supplemental Fig.
In summary, we find xenomiRs absent from most human tissue samples. In the samples where they are present, they are very lowly abundant. XenomiRs are not enriched in tissues exposed to dietary intake

Of the tissues in the human body, we would expect liver and blood cells to be relatively exposed to dietary molecules. Conversely, brain cells would be insulated from miRNA molecules by the BBB (Pardridge 2005; Boudreau et al. 2011). Despite this, we found that xenomiRs were present in similar fractions (28%) in the exposed liver and in the relatively protected brain samples. In addition, xenomiRs were present at a lower fraction of the blood cells (7%) than in the brain and liver samples (28%) (Supplemental Fig. 4: p1.1). Since the BBB can be permeabilized by neurodegenerative diseases, we repeated the analyses above considering only healthy control brains, with the comparable results (Supplemental Fig. 4: p1.2). These results support a nondietary origin of the detected xenomiRs. Interestingly, summed over all the 392 tissue samples, the xenomiRs were significantly overrepresented in older studies from before the year 2013 relative to the newer studies (88% versus 12%, P-value <0.001, Supplemental Fig. 4: p2). In addition, the older studies were more likely to contain xenomiRs from multiple clades than were the recent ones (P-value <0.001, Supplemental Fig. 4: p3). This observation suggests that the presence of xenomiRs in tissue samples is influenced by technical factors.

XenomiRs are commonly present in body fluids

We next analyzed the occurrence of miRNAs in the 432 human body fluid data sets, comprised of samples from serum, plasma, exosome, and CSF. As in the tissue samples, we found primate-specific (presumably endogenous human) miRNAs present in all body fluid samples (Fig. 3). However, the primate-specific miRNAs were overall less abundant and less diverse (in terms of distinct miRNA families) in the body fluid samples than in the tissue samples described above (median of count: 204 versus 979, P-value <0.001, distinct miRNA families: 11 versus 17, P-value <0.001; Supplemental Fig. 4: p4–5). Interestingly, xenomiRs were present in more body fluid samples (69% of samples) than in tissue samples.
The most common xenomiRs were from rodent, dicot, and insect clades (Supplemental Table 3B). We found that the rodent, euphyllophyta, lophotrochozoa, and insect xenomiRs group by study in body fluid samples (permutation test P-value <0.05) (Supplemental Fig. 4: p7); for instance, serum and CSF samples from study PRJNA241469 contained numerous rodent sequences, while serum samples from study GSE59565 had many insect sequences. The body fluids had low abundance and diversity of xenomiRs, with a median five xenomiR read counts and two distinct xenomiR families. These abundances and diversities were comparable in the tissue samples (P-value = 0.278 and P-value = 0.870, respectively, Supplemental Fig. 4: p8–9). Interestingly, xenomiRs were found in comparable fractions of serum (64%) and CSF (70%) from the study PRJNA241469, even though the CSF is separated from the bloodstream by the BBB (P-value = 0.312, Supplemental Fig. 4: p10). Serum and CSF xenomiR levels were also similar when considering only CSF from healthy individuals (χ² test, P-value = 0.476). To demonstrate that our findings are not biased by the choice of miRNA database, we repeated the analysis using only the highly curated miRNAs from MirGeneDB (Fromm et al. 2015). For zebrafish, chicken, and mouse we find exactly the same amount of miRNAs reported using miRBase and MirGeneDB (Supplemental Fig. 5). For human, we saw an imperfect correlation (Pearson’s correlation coefficient = 0.61) between miRBase and MirGeneDB. However, all our conclusions regarding P-values and grouping of samples still hold using the miRNAs reported by MirGeneDB (Supplemental Fig. 5). Overall, these results show that xenomiRs are present, although at very low levels, in most of the studied body fluid samples.

Most xenomiRs originate from unlikely food sources

If xenomiRs have dietary origins, we would expect that their composition would reflect the common human diet. For instance, based on our dietary preferences, one can expect a strong presence of crop plants (dicots, monocots, and euphyllophyta), fish and birds and a low presence of animal sponges, nematodes, insects, lophotrochozoa, echinoderms, and rodents, as these animals rarely constitute the human diet. Surprisingly, we find that rodents and dicots are the two clades that have the strongest xenomiR presence in the 432 body fluid samples (each is present in 34% and 29% of the samples, respectively) (Fig. 4; Supplemental Fig. 3B). Insect and euphyllophyta xenomiRs are also common (present in 13% and 11% of the samples). Even though birds and fish both comprise common human food sources, these
xenomiRs were only rarely observed in the serum, plasma, and exosome samples (0.8% and 4% percentages of samples, respectively). Regarding the prevalence of rodent sequences in the body fluid samples, rodents are accepted as a source of protein in the tropical world, but even there they only served as a supplement to the regular diets of humans (Fiedler 1990). Notably, rodents are of course widely used as model organisms (Wolff and Sherman 2008). Since most rodent-specific xenomiRs originate from a single study PRJNA241469 where the subjects were recruited in the USA, we speculate that the rodent xenomiRs are more likely derived from model organisms such as mouse and rat rather than from rodents supplied as a meat source. Regarding the insect sequences, it is not surprising that the human diet comprises traces of insects even in the developed world, as for instance wheat flour can harbor more than 150 insect fragments per 100 g (U.S. Food and Drug Administration). However, the human diet certainly includes many orders of magnitude more plant than insect material, and this disproportion is not reflected in the xenomiR abundances. We observe that although the abundances of plant dicot xenomiRs were statistically significantly higher than the insect (count median: three versus two, \( P\)-value = 0.004, Supplemental Fig. 4: p12), the count difference was not as huge as the disproportion of plant–insect content in diet. Last, the euphyllophyta xenomiRs like the previously reported MIR168 were present in a minority (11%) of the body fluid samples, showing that it is not commonly found in humans. These results show that the xenomiR composition does not reflect human food sources.

In summary, the 824 data sets comprise a total of 10.0 billion sequencing reads. We found that 3.4 billion (33%) reads are human miRNAs, of which 1.3 million reads (0.04% of human miRNAs) are primate-specific. The primate-specific miRNA reads are substantially less than the total number of human miRNAs in the samples, since the primate-specific miRNAs tend to be lowly expressed. Generally speaking, xenomiRs are lowly abundant, comprising only 0.001% (40,055 reads) of human miRNA reads (Fig. 4; Supplemental Table 3).

Human miRNA abundances are determined by body fluid type

Sequencing results are known to be influenced by technical factors that are study-specific (“batch effects”) (Akey et al. 2007; Leek 2014). To investigate if the measured human miRNA composition of the body fluid samples is more strongly influenced by biology or batch effects, we grouped the samples by principal component analysis (PCA). Since different tissues and cell types are known to have distinct miRNA profiles (Landgraf et al. 2007), we expected the body fluid samples to group according to their type (serum, exosome, etc.). Interestingly, we found that samples group both according to study (Fig. 5A) and also according to their sample type (Fig. 5B). However, hierarchical clustering revealed that the samples group primarily by sample type and secondarily by study (Supplemental Fig. 6). This is also observed in the PCA analysis where the samples grouped according to study, but only within the region primarily defined by the sample type (Fig. 5A,B). For instance, the serum samples from study PRJNA241469 clearly grouped with the other serum types, rather than with the CSF samples from the same study. These results show that the measured endogenous miRNA composition is determined primarily by the body fluid type.

XenomiR compositions are determined by technical batch effects

We next studied the patterns of the body fluid samples when grouped by xenomiR composition. If xenomiRs originate from diet, we would expect the samples to form clear dietary groups independent of the studies. For instance, samples...
from individuals with a vegetarian diet would group together, and similarly with individuals who had consumed fish or chicken. In contrast, we observed that the samples clearly grouped according to studies, suggesting technical batch effects (Fig. 5C, permutation test on hierarchical clustering showed four out of eight studies were significantly grouped, \(P\)-value <0.05, Supplemental Fig. 4: p13). This is in agreement with the hierarchical clustering analysis (Supplemental Fig. 7). This batch effect was particularly pronounced for studies PRJNA241469, GSE45722, GSE58410, and GSE59565 that had not been conducted with dietary restrictions. Further, we found four features that strongly determine the grouping: the presence of dicot, insect, lophotrochozoa, and rodent xenomiRs. As mentioned before, of these only the dicot xenomiRs are likely to have dietary origins. We did not observe any tendency for the samples to group according to the body fluid type when sorted by xenomiRs (Fig. 5D). These results indicate that xenomiR composition in human body fluids is determined by technical rather than dietary factors.

**Interindividual analyses of serum and CSF defy dietary origination of xenomiRs**

If xenomiRs had dietary origin, we would expect individuals with xenomiRs in CSF to have the same xenomiRs in the serum, since the foreign transcripts would need to enter the brain through the blood circulatory system. In the one study where CSF and serum small RNAs were profiled from the same individuals, we tested whether there is a significant tendency for xenomiRs to co-occur in CSF and serum of the same individual. Using binomial statistics with a significant level at 0.01, we found no significant tendency for xenomiRs from the euphyllophyta plant (\(P\)-value = 0.16), dicot plant (\(P\)-value = 0.04), insect (\(P\)-value = 0.27), or rodent (\(P\)-value = 0.08) to co-occur in the CSF and serum from the same individual, using binomial statistics with a significant level at 0.01. These results do not support that xenomiRs detected in CSF samples have entered through the bloodstream, and thus do not support a dietary origin of xenomiRs.
Controlled feeding in rat does not support dietary transfer of xenoRNAs

To investigate the influence of diet on blood xenomiR abundances, we performed a controlled feeding study in adult rats. Three different diets were applied and for each diet group we used three biological replicates: (i) three rats were fed only monocot plant material (rice), (ii) three rats were fed only dicot material (potatoes), and (iii) three rats were fed husbandry chow containing a defined mixture of grains, cereals, vitamins, minerals, and fats. After 28 d of controlled feeding (see Materials and Methods), small RNAs were isolated from the serum and sequenced. In these sRNA-seq libraries, we profiled rat miRNAs and searched for xenomiRs using the conservative method that we previously applied to the human data (Materials and Methods). If xenomiRs are indeed dietary-dependent, we would expect to see monocot-specific miRNAs in the rice-fed rats, dicot-specific sequences in the potato-fed rats, and both types in the rats that were fed chow. As expected, we identified abundant rodent-specific miRNAs in all nine samples (Supplemental Fig. 8). Interestingly, we observed a lack of plant xenomiRs in the rat serum. Specifically, we only observed two plant euphyllophyte sequences in a single rat sample.

We applied even more sensitive computational methods, by considering not only clade-specific sequences, but also miRNAs that are conserved but differ in sequence between species (see Materials and Methods). This method can potentially cause false positives when miRNAs are edited or subjected to sequencing errors. To account for this, we included negative and positive control samples. We used public Arabidopsis sRNA-seq data as negative controls and public rice and potato sRNA-seq data as positive controls. If our computational approach is sound, we would expect many of the rice and potato sequences to be flagged as belonging to their respective species, but few of the Arabidopsis sequences to be flagged as rice or potato. Indeed the rice and potato positive control sRNAs were clearly identified as such, each with >20,000 reads, while the relatively few Arabidopsis sequences misidentified as rice suggest relatively low false-positive rates (Fig. 6). As in the previous analysis, abundant rat miRNAs were observed (>10,000 reads summing over triplicates); however, only three rice-specific reads were seen in the rice-fed rats, and two rice-specific reads in the potato-fed rats. Given these extremely low numbers, and the fact that we observe rice-specific reads in the potato-fed rats, these are likely false positives of our sensitive computational method.

Controlled feeding in piglets does not support transfer of milk xenomiRs

It has been suggested that miRNAs can be transferred from breast milk to the bloodstream (Munch et al. 2013; Baier et al. 2014; Izumi et al. 2015), although these results have also been contested (Laubier et al. 2015; Title et al. 2015).

DISCUSSION

Public sequence repositories represent valuable resources in these times, when generating sequencing data is easier than providing meaningful analyses of the data. We have here analyzed 824 public human data sets for the presence of xenomiRs and have applied numerous analyses to evaluate if xenomiRs originate from diet or contamination. The data sets cover most existing sRNA-seq samples from human body fluids. We find that body fluids contain more xenomiRs than tissues, consistent with a dietary origin. However, we find several lines of evidence that xenomiRs in human samples originate from contamination: (i) XenomiRs are not enriched in tissues that are exposed to dietary intake; (ii) most xenomiRs originate from unlikely food sources like rodent and insect; (iii) xenomiR composition group by study,
nous transcripts, depending on the miRNA binding strength at around 1000 to 10,000 copies per cell to regulate endogenous which can reach 50,000 copies per cell (Lappalainen et al. 2014). These low counts contrast those of individual endogenous miRNAs, miRNA copies per cell (Materials and Methods). These low amounts of contaminants can be amplified through additional cycles of PCR this can be compensated for the small amounts of input material, in some cases down to 10 ng or 100 µL of input fluids for the RNA extraction (Wang et al. 2012, 2013), and this can be compensated through additional cycles of PCR amplification (Wang et al. 2012). This potentially means that very small amounts of contaminants can be amplified to yield detectable sequence read output.

Interestingly, we do observe a significant difference between the content of xenomiRs in tissues (present in 17% of samples) versus body fluids (present in 69% of samples). We considered possible technical sources of this difference. The standard Illumina protocol for sRNA-seq library preparation requires relatively large amounts of RNA (>1 µg of total RNA). This is easily obtained with tissue samples, but it can be difficult to extract these amounts of RNA from body fluids (Burgos et al. 2013). Therefore, sequencing of small RNAs from body fluids often use lower amounts of input material, in some cases down to 10 ng or 100 µL of input fluids for the RNA extraction (Wang et al. 2012, 2013), and this can be compensated through additional cycles of PCR amplification (Wang et al. 2012). This potentially means that very small amounts of contaminants can be amplified to yield detectable sequence read output.

Although xenomiRs were observed in most body fluid samples, we estimate that the levels are very low, at around 5 miRNA copies per cell (Materials and Methods). These low counts contrast those of individual endogenous miRNAs, which can reach 50,000 copies per cell (Lappalainen et al. 2013). It has been estimated that miRNAs need to be present at around 1000 to 10,000 copies per cell to regulate endogenous transcripts, depending on the miRNA binding strength and prevalence of targets (Denzler et al. 2014). Thus, even if the xenomiRs would be of dietary origin, they would be unlikely to be sufficiently abundant to affect transcripts in the bloodstream. On the other hand, given that the evidence suggests that xenomiRs are contaminants, they are represented by so few sequencing reads that they are unlikely to perturb any computational analyses. Thus, the xenomiRs likely represent trace contaminations that will have no consequences on the conclusions of the studies.

We realize that the numbers of clade-specific miRNAs differ from clade to clade. This reflects incomplete miRNA annotations, but also that some clades have gained more miRNAs than others in evolution (Sempere et al. 2006; Wheeler et al. 2009; Taylor et al. 2014). However, this is unlikely to confound our overall results. In particular, we have good coverage of the plant miRNAs, including MIR168, so the relative paucity of plant sequences in human biofluids cannot be attributed to our annotations. Similarly, we realize that our investigation of xenomiRs in body liquids is biased toward one comprehensive study that included more than half of our serum and CSF samples (study PRJNA241469). However, our main conclusions hold also when these samples are disregarded. Specifically, unlikely animal food sources such as insect and lophotrochozoa still dominate, while fish and bird xenomiRs are rare. Importantly, the samples still group according to the studies, suggesting batch effects. This shows that our conclusions are robust to the exclusion of particular studies.

In summary, our data support a technical rather than dietary origin of human xenomiRs. We have made the processed sequences from the data sets available, so that other researchers can extend our research. In addition, we hope that the >800 data sets, processed uniformly with a standardized pipeline, can function as a resource for the wider research community, for instance in the field of biomarker discovery.

FIGURE 7. Expression of cow and maize miRNAs. Sequenced RNAs that unambiguously mapped to cow and maize miRNAs were identified and summed up. The samples include sRNAs from horse (negative control), from pigs fed maize and pigs fed cow milk, and from cow and maize (positive control). The number above each bar indicates the number of sequenced RNAs mapping to cow or maize miRNAs.

Meta-study reveals xenomiRs are likely artifacts

Consistent with batch effects; and (iv) there is no co-occurrence of xenomiRs in CSF and serum from the same human individuals. In addition, we find no evidence of dietary transfer of xenomiRs in our controlled feeding studies in rat and piglet. These results lead us to conclude that dietary transfer of miRNAs into human tissues or body fluids are unlikely to be prevalent in humans.

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MATERIALS AND METHODS

Sequencing data collection

GSE accession numbers were collected from NCBI GEO database using the searching strategy "("micrornas[MeSH Terms] OR miRNA[All Fields]) AND "Non-coding RNA profiling by high-throughput sequencing"[All Fields]) AND "Homo sapiens"[porgn]" (271 GSE IDs, 5958 sequencing data sets, 12/08/2015). According to the GSE IDs, the metadata of contained data sets (XML format) were queried using esearch and efetch from Entrez Direct (Kans 2011). Using bash regular expression pattern, 823 samples (from 49 GSE IDs) that originate from human blood, liver, and brain and sequenced on an Illumina platform were selected and downloaded from the Sequence Read Archive (SRA) FTP site. It is important to note that SRA data sets in this case could be any type of sequencing data such as sRNA-seq, RNA-seq, ChIP-seq, etc. An extra 395 sRNA-seq data sets on serum and CSF samples
with BioProject ID PRJNA241469 (Burgos et al. 2014) were downloaded from NCBI dbGaP. In total, 1218 data sets were collected.

De novo 3′ adapter identification
The SRA format data sets were converted to FASTQ files using fastq-dump with option ‘-split-3’ from SRA Toolkit v 2.3.5. The paired-end reads were split into two FASTQ files with corresponding suffix _1.fastq and _2.fastq, while single reads were extracted into one FASTQ file. According to the file name, 75 paired-end sequencing data sets were removed. Since 3′ adapter sequences were not always provided together with FASTQ files and trimming the right adapter sequence is essential for the later efficient miRNA identification, all the remaining FASTQ files were processed for 3′ adapter identification using our in-house script in the following manner: Each FASTQ file was converted to FASTQ file with collapsing duplicate reads to speed up later processing. Homo sapiens mature miRNA sequences from miRBase v 21 were mapped to the reads of each FASTQ file considering all the perfect alignment using bowtie v 1.1.0. The 8 nucleotides (nt) subsequent to the aligned location in the FASTA reads were recorded and the most frequently appearing 8-mer was assigned as potential 3′ adapter sequence for that file. Besides determining the 3′ adapter sequences, the most abundant 8-mer can also approximately represent the number of miRNA reads profiled in that file.

Selection of sRNA-seq data
To exclude other types of sequencing data such as RNA-seq data, the abundance and sequences of the identified 3′ adapter were used as two filtering parameters. First, the file that has less than 100,000 identified 3′ adapter sequences (artificial cutoff) was excluded. The other types of libraries are not designed to capture miRNAs and unlikely to have a large amount of miRNA reads and the commonly used 3′ adapter sequence for sRNA-seq. Although some sRNA-seq data sets with low miRNA content might be discarded as well, these data sets are less likely to ensure a good representation of the eukaryotic small transcriptome. Second, among all the identified 3′ adapter sequences, only the data sets (n = 824) with the commonly used sRNA-seq 3′ adapter sequences (first 8-mer: TGGAAATTCT, TCCTGATGCG, ATCTCGTAT, AGATCGGA) were selected, as they are most likely to be sRNA-seq data (Supplemental Table 1).

sRNA-seq quality control
The raw files of these sRNA-seq data sets were processed applying the same quality control pipeline. The low quality, low complexity reads were removed using FASTX-Toolkit and custom script. The remaining reads were processed to check whether they contained the 3′ adapter sequence (8-mer) using regular expression of the Python programming language. For each adapter recognized read, the last appearing 8-mer together with subsequent nucleotides was trimmed. Reads shorter than 18 nt after adapter removal were discarded, since they are too short to be unambiguously traced back to reference sequences. The reads passing these quality control filters were collapsed to a FASTA file (Supplemental source) where the number after “_x” in the ID field indicates the read abundance. The FASTA files were used for miRNA expression analysis and clade-specific miRNA identification.

miRNA expression analysis
miRNA expression analysis was performed using Miraligner from Seqbuster (Pantano et al. 2010) with the Homo sapiens precursor miRNA (pre-miRNA) sequences and mature miRNA locations from miRBase v18 as reference. Briefly, the collapsed reads of each FASTA file were mapped to pre-miRNA sequences with allowing one mismatch (miraligner option ‘-sub 1’). Pre-miRNA and mature miRNA can be modified by extension, trimming, and substitution of nucleotides, resulting in distinct mature miRNA isoforms that have variations with respect to the reference sequences from miRBase (Ameres and Zamore 2013). The reads were also allowed to trim three nucleotides (-trim 3) or add one additional nucleotide (-add 1) in either the 3′ or 5′ end to reach the highest amount of alignments to pre-miRNA sequences. The miRNA expression matrix of 824 samples × 1787 known miRNAs (Supplemental Table 4) was generated by counting the number of reads that aligned to each annotated mature miRNA location on pre-miRNA.

Clade-specific miRNA identification and counting
According to the previous curated clade-specific miRNA family numbers (Supplemental Table 2), the reference catalog of clade-specific miRNA sequences was extracted from miRBase v21 by selecting the mature miRNAs with ID containing the family numbers using regular expression of the Perl programming language. For instance, the primate 1200 family yields two sequences with miRBase ID hsa-miR-1200 and ppy-miR-1200. A read was identified as a clade-specific miRNA read and assigned to the corresponding family such as primate_1200 if its first 20 nt are matched to the first 20 nt of the reference sequences. By counting how many reads were assigned to each family, we obtained a family matrix, including 129 clade-specific miRNA families from 16 clades × 824 samples (Supplemental Table 5). By further summing up the counts from the same clade, we got a clade matrix, including 16 clades × 824 samples (Supplemental Table 6). Note that we discarded the nematode miRNAs that were detected in samples from Bioproject PRJNA241469, where the nematode miRNAs were spiked in on purpose to normalize the human miRNA abundance. The two count matrices were used to generate Figures 2, 3. A summary table showing the total number of raw reads, human miRNA, primate-specific miRNA, and xenomiRNA clade-specific miRNA counts can be found in Supplemental Table 3.

Statistical analysis
The associations between the presence of xenomiRs or primate miRNAs with biological origins, study year, or sample types were evaluated with likelihood-ratio χ² test using loglm() function from MASS package in R v 3.1.1 (R Development Core Team 2010). If the χ² test revealed a strong relationship between multiple variables (n > 2) and outcomes, the partitioning analysis was further used to resolve which variable is associated. Since xenomiR counts were continuous variables and skewed distributed, we applied the Wilcoxon rank sum test for the comparisons that involved xenomiR abundance in different conditions; for instance, if xenomiR counts are different in body fluid versus tissue samples, we use wilcox.test() function in R. To test whether the hierarchically clustered samples are randomly distributed and whether the xenomiRs from each clade are
randomly present across samples, we applied a permutation test, where the sample IDs were randomly shuffled 1000 times to get the sampling distribution of test statistics. The ranking of the real test statistic among the shuffled test statistic gives a P-value. A small P-value is assigned if the real test statistic is beyond or located to the tails of the distribution.

Principal component analysis

The counts in the family matrix (Supplemental Table 5) and the clade matrix (Supplemental Table 6) were first normalized to reads per million (RPM) for each sample.

$$n_{rs} = c_{rs}/c_i \times 10^6,$$

where $c_{rs}$ is the raw counts of clade-specific family $r$ (ranged from 1 to 129) or clade $r$ (ranged from 1 to 16) for sample $s$, $c_i$ represents the total number of endogenous human miRNA counts estimated by seqbuster (see section “miRNA Expression Analysis”) for sample $s$. $n_{rs}$ is RPM of clade-specific family or clade $r$ that normalized to endogenous miRNA counts for sample $s$. In order to get a clear pattern of sample grouping, the $n_{rs}$ is further logarithm-transformed to $\log_2(n_{rs}+1)$.

The principal component analysis (PCA) of primate-specific miRNA families was performed on a subset of the normalized matrix including 41 primate-specific miRNA families $\times$ 432 body fluid samples, using the prcomp() function in R v 3.1.1. The sample points were projected in a subspace formed by the first and second principal components that explained 27.3% of the variability. The sample points were colored based on the study (Fig. 5A) and the principal components that explained 27.3% of the variability. The sample points were colored based on the study (Fig. 5A) and the principal component analysis of primate-specific miRNA families was much sparser than the corresponding primate matrix, containing many zero values. Therefore we summed up the xenomiR family counts of families belonging to the same clade, where the sample IDs were randomly shuffled 1000 times to get the sampling distribution of test statistics. The ranking of the real test statistic among the shuffled test statistic gives a P-value. A small P-value is assigned if the real test statistic is beyond or located to the tails of the distribution.

Feeding study library preparation and sRNA sequencing

Extracted total serum RNA from the two different animal models was concentrated in a speedvac to a final volume of 10 µL. From this 5µL was used as the input in the libraries, which were prepared accordingly with TruSeq Small RNA Sample Preparation kit (Illumina). We made one modification to the Illumina protocol, since we increased the number of PCR cycles from 11 to 16 for total RNA with a 1 µg as the starting material. Additionally, we applied TruSeq SR Cluster Kit v3 for cluster generation, and furthermore, for sequencing, TruSeq SBS Kit v3 was applied. Both were used according to the instructions. sRNA sequencing was conducted on an Illumina HiSeq2000 instrument at the core facility at the Centre for Genomic Regulation (CRG). We sequenced libraries as 50-bp single-end with a total fragment size of ~150 bp reads. Base calling and quality score calculation was performed from raw intensities using Illumina’s pipeline version 1.8.1.

Feeding study sRNA preprocessing and analysis

The sensitive sequence analyses applied to the feeding study data were as follows. The called rat and pig reads were trimmed with the command line: fastx_trimmer -f 1 -l 36 and low-quality reads discarded with fastx_artifacts_filter using the options -q 10 -p 50 (http://hannonlab.cshl.edu/fastx_toolkit/). Adapters were clipped using the AdRec.jar program from the Seqbuster suite with the following options: java -jar AdRec.jar 1 80 3. A custom search subsequently clipped shorter adapters: If there were no matches to the first 8 nt, then matches to the first 7 nt of the adapter were searched in the last 7 nt of the read, then matches of the first six to the last six positions and so on. Reads that had no matches were retained, but not clipped. Last, reads shorter than 18 nt were discarded. The nine rat data sets were analyzed with public Arabidopsis thaliana
Kang et al.

Estimating miRNA copies per cell

We assume that a mammalian cell contains around 100,000 endogenous miRNA molecules per cell (Calabrese et al. 2007). We then estimate the number of copies of an individual miRNA as: TPM (transcripts per million) \times \frac{100,000}{1,000,000}.

DATA DEPOSITION

Raw reads in the form of fastq files from the animal feeding studies have been stored in Gene Expression Omnibus GEO under accession number GSE92897. Furthermore, the processed sequence data from human tissue and body fluid in the form of fasta files can be downloaded from https://figshare.com/s/36e1c536a573011e4248.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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