gga-miRNOME, a microRNA-sequencing dataset from chick embryonic tissues

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MicroRNAs (miRNAs) are small non-coding RNA molecules, with sizes ranging from 18 to 25 nucleotides, which are key players in gene expression regulation. These molecules play an important role in fine-tuning early vertebrate embryo development. However, there are scarce publicly available miRNA datasets from non-mammal embryos, such as the chicken (Gallus gallus), which is a classical model system to study vertebrate embryogenesis. Here, we performed microRNA-sequencing to characterize the early stages of trunk and limb development in the chick embryo. For this, we profiled three chick embryonic tissues, namely, Undetermined Presomitic Mesoderm (PSM_U), Determined Presomitic Mesoderm (PSM_D) and Forelimb Distal Cyclic Domain (DCD). We identified 926 known miRNAs, and 1,141 novel candidate miRNAs, which nearly duplicates the number of Gallus gallus entries in the miRBase database. These data will greatly benefit the avian research community, particularly by highlighting new miRNAs potentially involved in the regulation of early vertebrate embryo development, that can be prioritized for further experimental testing.

Background & Summary

MicroRNAs (miRNAs) are small, single-stranded RNAs with sizes ranging from 18 to 25 nucleotides that are involved in gene expression regulation. This is achieved via post-transcriptional silencing of complementary messenger RNA (mRNA) targets by repression of translation and/or mRNA degradation¹. miRNAs were initially called small temporal RNAs (stRNAs), since they were first described as essential for proper developmental stage transition in the C. elegans life cycle². Today they are recognized to act as gatekeepers of developmental time in many other systems, by mediating cell proliferation-to-differentiation transitions³.

The canonical pathway of miRNA biogenesis starts with the transcription of a primary miRNA (pri-miRNA) by RNA polymerase II. The pri-miRNA forms a hairpin that is recognized by DGCR8, which recruits a Class 2 ribonuclease III enzyme, Drosha. This enzyme cleaves the RNA releasing the hairpin, called precursor miRNA (pre-miRNA), which is then exported to the cytoplasm via the Exportin-5 transporter. Here, it is recognized by a second Class 2 ribonuclease III enzyme, Dicer, that cleaves the loop from the hairpin releasing a small double-stranded RNA. One of the strands binds to an Argonaute protein from the RNA-induced silencing complex (RISC), while the other is degraded. At this point, the mature miRNA selectively recognizes and binds to the 3’ untranslated region (3’UTR) of its target mRNA through a small 2–7 nucleotide seed region, leading to RISC-mediated mRNA degradation and/or translational repression¹.

An essential step in addressing miRNA-mediated regulation of gene expression is to identify and quantify the miRNAs present in the biological system of interest. High throughput miRNA profiling studies have identified thousands of miRNAs in human and mouse samples⁴. However, this effort has been lagging behind in other model organisms, hindering the elucidation of their role in these systems. This is the case of the chicken (Gallus gallus) embryo, a well-established model for studying human embryogenesis due to its extraordinary molecular and morphological similarities in the early stages of development, alongside the ease of experimental

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manipulation it offers. It was in the chicken embryo that the molecular embryonic clock (EC) underlying the periodic formation of vertebral precursors was first described. EC genes present cyclic expression maintained by negative feedback regulation in the posterior undetermined presomitic mesoderm (PSM), which gradually slows in the anterior PSM and halts in the segmented somites. The periodicity of gene expression oscillations in the PSM is species-specific but can also differ in different tissues of the same organism. Namely, *hairy*2 gene expression oscillates with a periodicity of 90 min in the chick PSM and 6 h in the distal cyclic domain (DCD) of the developing forelimb bud.

mRNA instability is essential for EC cycles of expression and there is evidence of a miRNA-dependent regulation of EC gene oscillations. Namely, miR-125a-5p is required for cyclic *LFGN* expression in the chick PSM and miR-9 drives *Hes1* oscillations in mouse neural progenitor cells. Additionally, we previously showed that the genes encoding the enzymatic machinery for miRNA biogenesis are expressed in both the chick PSM and forelimb bud, tissues where the EC is oscillating.

A thorough characterization of the role of small RNAs in chick embryo development and in the regulation of the EC has been hampered by the scarcity of miRNA expression datasets in embryonic tissues of this model system. To overcome this limitation, we performed a miRNA profiling analysis (miRNA-Seq) of three different tissues of the developing chick embryo (Fig. 1a,b). Namely, two regions of the PSM - Undetermined Presomitic Mesoderm (PSM_U) and Determined Presomitic Mesoderm (PSM_D) - and the Forelimb Distal Cyclic Domain (Limb). We report the identification of 926 known miRNAs, and 1,141 candidate novel miRNAs, not previously described in chicken. Accordingly, we believe that this will be an invaluable data resource for the research community studying miRNA-mediated gene expression in early vertebrate development, particularly in the chick embryo.

**Methods**

**Embryos.** Fertilized *Gallus gallus* eggs (Pintobar, Portugal) were incubated at 38 °C in a humidified atmosphere for two or four days to obtain embryos in stages HH12–13 and HH20–22, respectively.

**Sample collection.** Presomitic mesoderm (PSM) tissues were isolated from embryos in stages HH12–13. To obtain these samples, embryos were collected from 48-h-incubated eggs, placed in a petri dish containing phosphate buffer saline (PBS) solution and staged according to Hamburger and Hamilton. Only the embryos in stages HH12–13 were selected for further use. The embryos were then placed ventral side up in PBS and 4 μL of Pancreatin (25 mg/mL) (Sigma #8049-47-6) was added to the surface of the embryo. After 3 to 5 minutes, pancreatin was inactivated with goat serum (Gibco #16210-072). The mesoderm located on either side of the neural tube was isolated from all surrounding tissues and divided into determined PSM (upper one-third portion) and undetermined PSM (caudal two-thirds) (Fig. 1a,b). Due to the extraordinarily small size of these tissues, 20 pairs of PSM portions were pooled together for RNA extraction from each biological sample. The samples were snap frozen in liquid nitrogen and stored at -80 °C.

Distal Cyclic Domain (DCD Limb) tissues were isolated from embryos at stages HH20–22 (Fig. 1a,b). Embryos were collected from fertilized eggs incubated for four days, placed in PBS and staged according to Hamburger and Hamilton. Only the embryos in stages HH20–22 were selected for further use. The limb tissue (distal medial portion of the forelimb bud) was manually dissected using forceps. 20 DCD Limb pairs were pooled together for each sample, snap frozen in liquid nitrogen and stored at -80 °C.

**RNA extraction.** Biological samples were defrosted on ice. Total RNA was extracted using TRIzol Reagent (Invitrogen #15596-018) according to the manufacturer’s instructions with slight adaptations, namely, the aqueous phase from the first step of extraction was washed once with Phenol:Chloroform (Sigma #P2069) and then with Chloroform:isoamyl alcohol (24:1). The aqueous phase was recovered using Phase Lock Gel Heavy (5Prime #2302830) and RNA was precipitated by addition of 1/10 volume of 3 M sodium acetate, 2.5 volumes of 100% ethanol and 3 μL per mL of Linear Acrylamide (Ambion #AM9520). After one hour at -80 °C, the RNA was precipitated by centrifugation at 14,000 rpm for 30 minutes at 4 °C. The pellet was washed with 70% ethanol and centrifuged for 15 minutes at 4 °C, briefly air-dried and resuspended in 50 μL of MilliQ (Merck Millipore) purified water. The samples were quantified using NanoDrop 2000 (Thermo Scientific) and stored at -80 °C.

**RNA quality control.** A first-round of quality control was performed by Reverse Transcription-PCR. 100 ng of RNA was reverse transcribed using iScript™ cDNA Synthesis Kit (BioRad #1708890). Subsequent PCR for GAPDH was done using DreamTaq DNA Polymerase (Thermo Scientific™ #EP0701). In a second instance, RNA quality control was performed using Experion™ RNA StdSens Analysis Kit (BioRad #700-7103) (Table 1). Only samples with an RQI (RNA Quality Indicator) equal to or above 8.5 were sent for sequencing.

**Library preparation and miRNA-sequencing.** The sequencing libraries were prepared using the NEBNext Multiplex Small RNA Library Prep Set for Illumina (NEB #E7300S/L Version 5.0), starting with 150 ng of total RNA as input. As a first step in the protocol, adaptors ligate directly to the small RNA fragments containing 5’ phosphate and 3’ OH, followed by cDNA generation and PCR amplification. 15 cycles of amplification were performed using specific SR primers for Illumina index and primer index of choice for each sample (according to NEB #E7300S/L Version 5.0 protocol).

Size distribution of the final library was assessed on Bioanalyzer (Agilent Technologies) with a DNA High Sensitivity kit (Agilent Technologies #5067-4626), and concentration was measured with Qubit® DNA High Sensitivity kit (Life Technologies #Q32854) in Qubit® 2.0 Flurometer (Life Technologies). Individual libraries that passed the QC step were pooled equimolarly in a 9-plex, and final pool was purified with SPRI select beads at a 1.3x bead ratio (Beckman Coulter #B23519). Pool was loaded to a single lane of an Illumina HiSeq 2000
sequencing instrument (Illumina Inc.) at 6 pM concentration and was sequenced in 50 bp single-read mode\textsuperscript{15}. Library construction and sequencing were performed at EMBL’s GeneCore facility in Heidelberg, Germany.

**Quality control of sequencing reads.** Sequencing reads were firstly evaluated using FastQC (version 0.11.5)\textsuperscript{16} to verify the overall read quality of each sample. One library (PSM_U2), from undetermined PSM, did not pass the quality control step, mainly due to its small library size, leading to its removal from further analyses (Fig. 2a).

**Quantification and normalization of annotated miRNAs.** For the remaining 8 samples that passed the quality control, annotated microRNA read counts were obtained using the Chimira software (version 1.5)\textsuperscript{17}. Briefly, the pipeline implemented in Chimira for miRNA-seq analysis comprises the following steps: firstly, the sequences are cleaned, trimmed, and size selected to remove adapters and low quality microRNA reads. Next, the

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### Experimental design

| RNA extraction | Library preparation | Sequencing | Quality Control |
|----------------|---------------------|------------|-----------------|
| TRizol isolation | NEBNext Multiplex Small RNA Library Prep Set for Illumina | Illumina HiSeq2000 platform | FASTQ files (raw sequences) |

**Data analysis**

- Summary statistics
- Principal Component Analysis (PCA)
- Expression profile for annotated miRNAs
- Expression profile for novel miRNAs

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**Protocol & Data analysis**

| RNA extraction | Library preparation | Sequencing | Quality Control |
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## Figure 1

Experimental design, protocol overview, and data analysis workflow. (a,b) Overview of the experimental design, showing the sampling sites of the chick embryonic tissues collected. (c) Pipeline for annotated miRNA-seq data analysis and novel miRNA prediction. PSM_D: determined Presomitic Mesoderm (PSM); PSM_U: undetermined PSM; DCD: Limb Distal Cyclic Domain.
reads passing the previous filters are mapped to *Gallus gallus* hairpin sequences present in miRBase (release 22)\(^4\) using BLASTn\(^18\) allowing up to two mismatches. Finally, a count-based miRNA expression dataset is generated\(^19\) and normalized across all samples using DESeq2\(^20\). Further data validation, visualization, and statistical analyses were conducted using the normalized log2 expression data.

**Detection, quantification and normalization of novel miRNAs.** Detection of novel miRNAs was performed using the Mirnovo tool (v1.0)\(^21\), which is a machine learning algorithm that predicts novel miRNAs by analysing structural features of miRNA precursor hairpin sequences gathered directly from small RNA-Sequencing data. Briefly, the Mirnovo pipeline entails the following steps: (i) adapter removal followed by sequence de-duplication; (ii) the tallied sequences then enter a series of clustering steps, followed by cluster refinement to obtain consensus sequences; (iii) the prediction step identifies known and novel miRNAs; and (iv) the final step, aligns the consensus sequences from all miRNAs (known and novel) to the reference genome. This was done by selecting the most stable hairpins (scored by Delta G free energy) found in a 90-nucleotide window around the consensus sequences followed by genomic feature calculation\(^21\).

The specific parameters used for Mirnovo were: *Gallus gallus* input species, using the Universal prediction model (since there are no models specifically trained for chicken), length filter between 16 and 28 nucleotides, minimum read depth of 5, minimum variants 1, and initial clustering using an alignment identity threshold of 0.9 (vsearch-id parameter).

The candidate novel miRNAs were then quantified and normalized using Chimira\(^17\) with the Mirnovo extension. The analysis was performed as described above for the annotated miRNAs, with the difference that the custom hairpin FASTA files output from Mirnovo for each sample were uploaded together with the corresponding raw FASTQ files.

Data resulting from this identification and quantification (i.e. hairpin sequences, genomic location, and normalized counts) is freely available\(^22\).

**microRNA expression profiling.** Using customized R scripts (R version 3.6)\(^23\), we conducted quality control analysis, and briefly inspected the profile of annotated and novel microRNA expression in each embryo tissue. For this we used R packages for data visualization, namely, Tidyverse\(^24\), UpSetR\(^25\), Patchwork\(^26\), and plot3D\(^27\).
Data Records
All sequencing data has been deposited in the ArrayExpress data repository with accession number E-MTAB-8176. This dataset consists of 8 microRNA expression raw data files in fastq format. Detailed experimental procedures and data analysis are also available there.

Processed data (in tabular text format) containing the log2 normalized counts of the sequencing reads for annotated miRNAs has been deposited in Figshare. Similarly, the list of predicted novel miRNAs, with sequence, and log2 normalized counts is available in Figshare.

All the sequencing data and the normalized miRNA expression counts are open. The R code used for the exploratory data analysis and visualizations are also freely available for consultation in Figshare.

Technical Validation
Quality control of microRNA-Seq data. The quality control of the raw sequencing reads was performed using FastQC to assess overall read quality and flag potentially poor-quality samples. All samples except one, passed the QC metrics performed by FastQC. The poor-quality sample presented a variable PHRED score distribution across the read length (Fig. 2a), as well as a very low total number of reads (244,296 reads compared to 3 million average reads in the other samples), hinting that the sequencing step was faulty, possibly due to sample...
The top-20 most expressed miRNAs (Fig. 5a,c) are found in all three tissues, with roughly comparable distributions in both annotated and novel miRNAs. Additionally, the intersection plot (Fig. 5b,d) clearly shows that the majority of miRNAs (637 in known miRNAs and 849 in novel miRNAs) are found in all three tissues. Importantly, each tissue presents exclusive miRNAs, namely 71 in Limb, 35 in determined PSM, and 8 in undetermined PSM.

To validate the read normalization and quantification steps, we evaluated the read distribution before and after normalization, and briefly compared the miRNA expression profile across tissues.

The distribution of the total number of reads (Fig. 4a,c) shows that there are some differences between the replicates before read count normalization, particularly for the Determined PSM tissue in the known miRNAs set. This fact is most likely a reflection of the embryo pooling strategy that might be contributing asymmetrically to the total miRNA amount present in each sample. After normalization (Fig. 4b,d), the distributions become more balanced between replicas, and therefore amenable for further expression comparisons between tissues. As expected, the miRNA expression distribution for all three tissues is positively skewed (even after log2 transformation), showing a long tail to the right.

miRNA Expression profile in the different tissues. Looking at the miRNA expression profile in the three tissues helps with uncovering possible experimental errors; for example, large asymmetries in the diversity of miRNAs found for each tissue could indicate faulty sequencing, or a total overlap of miRNA identities between tissues could indicate mislabelling or inadequate experimental design.

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in undetermined PSM for known miRNAs (Fig. 5b); and 51 in Limb, 41 in determined PSM, and 7 in undetermined PSM for novel miRNAs (Fig. 5d), showing that each sample is sufficiently different from the others, allowing for proper differentiation between tissues.

Usage Notes

The bioinformatics analysis described here made use of freely available software tools commonly used by the research community (Fig. 1c). There are alternative miRNA-seq analysis pipelines equally applicable to the FASTQ reads from *Gallus gallus*22, for example, miRDeep230, QuickMIRSeq31, and sRNAnalyzer32. For a recently published miRNA-seq analysis protocol, see Potla et al.33, discussing available individual tools for each step: (i) quality control (adaptor trimming, read quality/length filtering); (ii) read mapping; (iii) annotation (using miRBase); (iv) quantification; and optionally (v) detection of novel miRNAs.

These data can equally be used to seek the complete small RNAome, using for example the recently developed platform coMpsRA that is reported to identify and quantify diverse RNA molecule types, including miRNA, piRNA, snRNA, snoRNA, tRNA, and circRNA34.

The miRNA expression data herein reported19 will be useful to study gene regulation in the early phases of vertebrate embryo development, for example by performing differential expression and target gene annotation analyses. Some considerations should be taken into account for downstream analyses. Namely, the RNA was extracted from pools of 20 dissected tissues meaning that each sample represents an heterogeneous mixture of individuals, whose variability is present in the data. This is even more relevant if we consider that oscillations of clock gene expression occur in the tissues analysed. Thus, care should be taken when using such static sample datasets to contrast tissues with dynamical gene expression. Additionally, some SNPs can potentially interfere with the successful mapping of some miRNA transcripts that might have been discarded, and therefore cause an underrepresentation of expression for those miRNAs. Finally, for differential expression studies comprising the PSM_U tissue, since this group comprises only two replicates, the comparison will have lower statistical power to detect small effect sizes. Accordingly, appropriate statistical techniques should be applied to deal with this limitation.

Since the chicken genome annotation is not yet up-to-par with the annotations from other vertebrate genomes, most chicken miRNAs deposited in databases are not yet experimentally validated, and their target genes are based mostly on chicken-specific computational predictions. This study opens the door for new findings specific for birds, and for validation of known vertebrate miRNAs and their respective target genes. Finally, the predicted novel miRNAs22 represent an invaluable resource for the avian research community looking to experimentally validate novel candidate miRNAs acting in early vertebrate development capable of regulating
Fig. 5 Expression profiling and overlap between the three tissues for annotated (a,b) and novel (c,d) miRNAs. (a,c) Top 20 highly expressed miRNAs per tissue. (b,d) Intersection between miRNAs expressed in each tissue.

their gene of interest. Additionally, these data coupled with transcriptomics data for the same tissues can help uncover potential regulatory modules active in early vertebrate embryogenesis.

Code availability
Technical validation and data visualization was performed in RStudio (Version 1.1.463)35, using R (version 3.6)23, and Bioconductor (version 3.9)36, with packages tidverse (version 1.3.1)24, UpSetR (version 1.4.0)25, patchwork (version 1.1.1)26, plot3D (1.3)27. The R code used for these analyses, in the form of an annotated R notebook, is freely available in Figshare16. Additional software tools used to analyse this miRNA-seq dataset were the following: FastQC (version 0.11.5)16, Chimira (version 1.5)17, and Mirnovo (version 1.0)21 as described in the Methods section.

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Author contributions
I.D. performed data analysis, managed, and archived the data, and drafted the manuscript. G.C. collected the samples, performed the RNA extraction, and drafted the manuscript. N.T.D.A. and V.B. acquired the data. R.P.A. conceived the study, designed the experiments, coordinated the project, and drafted the manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare no competing interests.

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