1	Single paternal Dexamethasone challenge programs offspring metabolism and reveals
2	multiple candidates in RNA-mediated inheritance
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4 Summary

5 Single traumatic events that elicit an exaggerated stress response can lead to the development 6 of neuropsychiatric conditions. Rodent studies suggested germline RNA as a mediator of 7 effects of chronic environmental exposures to the progeny. The effects of an acute paternal 8 stress exposure on the germline and their potential consequences on offspring remain to be 9 seen. We find that acute administration of an agonist for the stress-sensitive Glucocorticoid 10 receptor, using the common corticosteroid Dexamethasone, affects the RNA payload of 11 mature sperm as soon as 3 hours post exposure. It further impacts early embryonic 12 transcriptional trajectories, as determined by single embryo sequencing, and metabolism in 13 the offspring.

We show persistent regulation of tRNA fragments in sperm and descendant 2-cell-embryos, suggesting transmission from sperm to embryo. Lastly, we unravel environmentally induced alterations in sperm circRNAs, and their targets in the early embryo, highlighting this class as additional candidate in RNA-mediated inheritance of disease-risk.

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19 Introduction

20 Acute stress elicits a complex but well-studied cascade of neuroendocrine responses 21 regulated by the hypothalamic pituitary adrenal axis. It involves the release of neuropeptides 22 in the brain that induce the secretion of corticosteroid hormones from the adrenals. These 23 hormones in turn activate two types of corticosteroid receptors, glucocorticoid receptors (GRs) 24 and mineralocorticoid receptor (MRs). These receptors are widely expressed throughout the 25 body and regulate gene expression, thus enabling physiological and behavioral adjustments 26 in response to stress (de Kloet, Joëls and Holsboer, 2005). In vulnerable individuals, this 27 response is excessive and it can lead to long lasting maladaptive changes with consequences 28 for psychological and metabolic health (Daskalakis et al., 2012).

1 It is also known that parental experiences can compromise the health of their progeny both in 2 humans (Pembrey et al., 2006; Heijmans et al., 2008; Bowers and Yehuda, 2016) and in 3 animal models (Benyshek, Johnston and Martin, 2006; Roth et al., 2009; Jimenez-Chillaron 4 et al., 2009; Carone et al., 2010; Pentinat et al., 2010; Shankar et al., 2010; Franklin et al., 5 2010, 2011; Morgan and Bale, 2011; Weiss et al., 2011; Dietz et al., 2011; Vassoler et al., 6 2013; Fullston et al., 2013; Martínez et al., 2014; Gapp et al., 2014; Marco et al., 2014; Rodgers 7 et al., 2015; Sharma et al., 2015; Wu et al., 2016; Chen et al., 2016; Y. Y. Zhang et al., 2018). 8 Research on the underlying mechanism of such transmission has found changes in germline 9 epigenetic make-up, in particular DNA methylation, histone post translational modifications 10 (PTMs), histone positioning and RNA (Gapp and Bohacek, 2017). These epigenetic regulators 11 are responsive to the environment and have been implicated in a variety of environmentally 12 induced diseases (Jirtle and Skinner, 2007). Altered modifications must circumvent epigenetic 13 reprogramming events in zygote and, depending on the timing of exposure, during germline 14 development (Bohacek and Mansuy, 2017; Gapp and Bohacek, 2017). In the male germline, 15 RNA is excluded from reprogramming and therefore a promising candidate for 16 transgenerational information delivery (Gapp and Bohacek, 2017; Bohacek and 17 Rassoulzadegan, 2019). Several studies carried out in D. melanogaster and C. elegans 18 reported on transgenerational inheritance of induced traits and provided firm evidence for the 19 involvement of small RNAs in the mechanism of transmission (Ashe et al., 2012; Grentzinger 20 et al., 2012; Shirayama et al., 2012). In mammals, a causal implication in the transmission of 21 environmentally induced effects across generations has been demonstrated for sperm RNA 22 only (Gapp et al., 2014; Grandjean et al., 2015; Sharma et al., 2015; Chen et al., 2016). Such 23 RNA differs substantially from somatic RNA since it mainly consists of small RNA, 24 predominantly tRNA-derived small fragments (tsRNAs), but also miRNAs, piRNAs and 25 circRNAs, among others (Chen, Yan and Duan, 2016; Gapp and Bohacek, 2017; Bohacek and 26 Rassoulzadegan, 2019). CircRNAs, comprise a very stable class of RNA that has recently 27 been observed to be present in high amounts in testis but also to some extent in sperm (Barrett 28 and Salzman, 2016). Some have been shown to act as miRNA sponges, thereby competing

with mRNA targets, while also regulating the expression of their host genes (Barrett and Salzman, 2016). Hence, circRNAs have a strong potential for amplifying an inherited signal, which makes them exceptionally interesting candidates for epigenetic germline inheritance. To date, the involvement of circRNAs in soma-to-germline signalling has not yet been investigated.

6 tsRNAs and miRNAs are crucial regulators of early embryonic development and players in 7 non-genetic inheritance (Gapp et al., 2014; Grandjean et al., 2015; Rodgers et al., 2015; 8 Sharma et al., 2015; Chen et al., 2016; Benito et al., 2018; Tyebji, Hannan and Tonkin, 2020). 9 They have been reported to be acquired through exosomal uptake during epididymal transfer 10 from caput to cauda epididymis (16,44). This might explain their responsiveness to 11 environmental perturbations, despite mature sperm's presumably transcriptionally silent state 12 caused by tightly packed chromatin. Sperm RNA can indeed change in response to chronic 13 stress or by chronic treatments that mimic stress exposure, such as repeated injection of GR 14 agonists (Rodgers et al., 2013; Gapp et al., 2014; Short et al., 2016; Wu et al., 2016). In mice, 15 uptake of epididymosomal miRNA was sufficient to replicate a chronic stress induced effect on 16 stress response in the offspring (Chan et al., 2020). Surprisingly, acute stress has also 17 recently been shown to affect offspring weight and glucose metabolism in mice (Hoyer et al., 18 2013) and some of these effects were germline dependent (Bohacek, von Werdt and Mansuy, 19 2016). Together these related lines of evidence led us to hypothesize that acute GR activation 20 has an intergenerational effect on offspring phenotype and that the transmission potentially 21 implicates changes in the germline. The male germline cells - including mature sperm 22 (Kaufmann et al., 1992; Haeussler and Claus, 2007) and their surrounding Sertoli cells (Hazra 23 et al., 2014) - as well as the epididymal epithelial cells (Silva et al., 2010) express GRs that 24 mediate the effects of glucocorticoids on transcription. Dexamethasone (Dex), a specific GR 25 agonist, is known to directly activate GR in the rat epididymis (Silva et al., 2014). It is unknown 26 whether acute stress affects sperm RNA, and if so, whether uptake via epididymosomes is 27 involved in establishing germline changes that are relevant for offspring phenotypic alterations.

1 Here we investigate the impact of acute GR agonist administration on the germline RNA 2 payload including circRNAs, at various time points post administration and interrogate the fate 3 of altered sperm RNA. We further test germ-line dependency of transmitted metabolic effects 4 and dissect the underlying molecular trajectories during early embryonic development using 5 single cell sequencing of in vitro fertilization (IVF) derived embryos. Identifying a readout of 6 transgenerational risk load at the level of the paternal sperm epigenome could pave the way 7 for future studies aiming at a prevention of the transmission of the effects of acute GR 8 activation to the offspring.

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11 Results

12 Effects of acute Dex injection on the germline small RNA payload

13 Two reports have suggested that a single foot shock could elicit effects on offspring phenotype 14 (Hoyer et al., 2013; Bohacek, von Werdt and Mansuy, 2016). To examine potential epigenetic 15 mediators of such acute stressful impacts we investigated sperm RNA of males 2 weeks after 16 a single activation of the GR (Figure 1A). This timeline was chosen to mimic the timing at which 17 breeding occurred when effects on offspring had been observed in a previous study (Bohacek, 18 von Werdt and Mansuy, 2016). We injected the specific GR agonist Dex once intraperitoneally 19 into 8 adult males. This drug is in frequent clinical use, now also as an apparently effective 20 treatment for patients suffering from lower respiratory tract infection as a consequence of 21 Covid-19 virus (EU Clinical Trials Register, no date; 'Biggest COVID-19 trial tests repurposed 22 drugs first', 2020; Horby et al., 2020). A sperm population was harvested from each animal 23 and RNA was extracted for ultra-deep small RNA sequencing, resulting in 16 libraries 24 representing one injected male each (8 vehicle and 8 Dex-injected). Purity of the sperm 25 samples was confirmed by inspecting RNA size profiles generated on the bioanalyzer to be 26 absent of ribosomal RNA peaks, that would indicate contamination by somatic cells. Reaching 27 an average of 55.4 million sequencing reads while also using randomized adapters for 3' 28 ligation put us in a position to reduce PCR biases (Dard-Dascot et al., 2018) and accurately

1 quantify less abundant miRNAs that are by far outnumbered in sperm by other small RNAs 2 e.g. tsRNAs(Peng et al., 2012). Our data showed an average of 60% mappable reads across 3 all libraries, including 34% of multimappers. We detected an expected dominant prevalence of 4 reads mapping to tsRNAs and abundant miRNAs in all samples (Supplementary Fig. 1A). 5 Differential gene expression analysis, using DEseq2 (Love, Huber and Anders, 2014), revealed 6 that a single acute activation of GR receptors induced changes in tsRNAs, miRNAs and rRNAs 7 collected 14 days post injection (Figure 1B,C, FDR q<0.05, Table 1,2), as has been observed 8 in response to chronic environmental stress previously (Rodgers et al., 2013; Gapp et al., 9 2014). Interestingly, tsRNA-Gly-GCC, a tsRNA previously associated with the effects of 10 nutritional challenge (Sharma et al., 2015), was among the most strongly altered tsRNAs. We 11 further detected changes in ribosomal RNA-derived RNA (rsRNA) as has been observed in 12 studies investigating the impact of high fat and high sugar diet (Y. Zhang et al., 2018; Nätt et 13 al., 2019) (Figure 1D,E). It should be noted that our analysis only detects relative changes 14 within the total small RNA pool. Thus, we cannot exclude that the apparent increase in rRNA 15 is due to changes in other RNA subtypes. Accordingly, despite the suggested predictive value 16 of altered rsRNAs for fertility (Hua et al., 2019) we did not observe changes in sperm count. 17 fertilization rate or litter sizes in the Dex treated sperm and resulting offspring (Supplementary 18 Fig. 6).

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20 Some recent publications have suggested that sperm miRNAs and tsRNAs are acquired during 21 epididymal transit from caput to cauda (Sharma et al., 2015, 2018; Conine et al., 2018). 22 Further, it was shown that changes in sperm tsRNAs, induced by chronic nutritional-challenge, 23 are acquired by uptake of distinct sets of tsRNAs (Sharma et al., 2015). To examine whether 24 the changes observed 2 weeks post Dex injection were also apparent in caput sperm before 25 epididymal transit we decided to assess a selection of small RNAs in sperm harvested from 26 caput epididymis using q-PCR. Out of 7 selected small RNAs encompassing both tsRNAs and 27 miRNs we found 6 unaltered (Figure 1F) in line with the assumption that epididymal transit is 28 required to allow epididysomal uptake leading to altered RNA cargo in mature sperm. Yet

tsArg-CCT-2 was consistently altered in caput sperm (Figure 1F) indicating that this change is
 either induced at an earlier transcriptional level during spermatogenesis, or that certain small
 RNAs are taken up from exosomes in caput epididymis.

To test these two hypotheses, we examined a small set of RNAs in serum circulating exosomes. Indeed, we detected increased levels of tsArg-CCT-2, while other tsRNAs and miRNAs did not show altered serum exosome payload (Figure 1G). Thus, it appears that tsArg-CCT-2 is taken up by sperm from exosomes in the caput, but that other small RNA changes are not necessarily reflected in the payload of circulating exosomes.

9 While the necessity of epididymal transit to acquire changes represents one explanatory 10 framework for the absence of change in all but one selected small RNA in caput sperm, 11 alternative explanations should also be considered. Our results could also indicate that 12 changes observed in mature sperm 14 days post Dex injection represent a highly specific 13 snapshot in time, that relies on the affected sperm cells to be in a specific developmental stage 14 at the time of treatment. Cells entering into more mature stages of sperm differentiation at a 15 later point after Dex administration, such as the here assessed cells sampled from caput would 16 then no longer display said changes. Therefore we cannot conclusively establish a reliance on 17 epididymosomal uptake. To further dissect the dependence on epididymosomal uptake during 18 transit from caput to cauda epididymis from a different angle, we assessed the mature sperm 19 small RNA payload at two time points, 3 hours and 7 days following injection (Figure 2A). Cells 20 collected from cauda 7 days after injection have already exited testis, and have had time to 21 pass through the entire epididymal tract before collection. In contrast, cells collected 3 hours 22 post injection have most likely not passed through the corpus epididiymis, and already reside 23 in cauda epididymis at the time of injection where sperm resides up to 5 days (Meistrich, 1975; 24 Dadoune and Alfonsi, 1984). Importantly, spontaneous ejaculation regularly voids cauda 25 epididymis of sperm, even in the absence of a mating partner(Huber and Bronson, 1980), 26 excluding the retention of "old" mature sperm in cauda for prolonged periods of time. The cells 27 collected 7 days after exposure therefore represent a mixture of cells that might have already

resided in the cauda and those cells that indeed passed through the corpus epididymis, yet
 the spontaneous ejaculation ensures that the sample predominantly contains the latter.

3 The collected samples were confirmed for their purity and again processed separately to 4 represent sperm from one animal per library. The resulting libraries were analysed jointly as to 5 test for (1) effects of Dex injection independent of sampling time post injection (2) effects of 6 sampling time post injection independent of Dex treatment and (3) effects depending on both 7 Dex injection and the sampling time post injection (interaction). We report an average of 64% 8 mappable reads including 46% of multimappers and observe that tsRNAs were significantly 9 affected by sampling time post injection independent of treatment. This demonstrates the 10 fluctuation of tsRNAs over time in response to external signals such as injections, or potentially 11 due to uncontrollable external inputs from the animal husbandry (Figure 2C). Interaction 12 between treatment and time was statistically significant for 27 tsRNA mapping loci including 13 Gly-GCC-6-1. All affected tsRNAs are upregulated after 7 days. 26 tsRNAs of them are 14 unchanged after 3 hours and one tsRNA (Thr-TGT1-1) is downregulated after 3 hours 15 (Supplementary Table2 sheet 3, q<0.05). This finding is consistent with the dominating view 16 that tsRNAs are acquired during epididymal transit from caput to cauda epididymis. However, 17 most tsRNAs that showed a significant change in response to treatment after 7 days, but not 18 after 3 hours (interaction between treatment and time post injection, Supplementary Table 2 19 sheet 3, q<0.05) were not persistently altered in the dataset of 14 days after injection (Table 20 1). This indicates that on the one hand changes in sperm RNA are dynamic and many do not 21 persist for prolonged time. On the other hand, this suggests that potentially relevant small RNA 22 changes mostly require either sperm to reside in testis at the time of exposure, or rely on a 23 prolonged residency in the exposed organism. Interestingly, we also detected 2 exceptions 24 that show a significant group effect across 3 hours and 7 days. tsRNA-Leu-CAA and tsRNA-25 Arg-CCT (Figure 2B) were persistently affected 3 hours and 7 days post exposure, that 26 necessarily requires a mode of rapid acquisition of tsRNA-changes in cauda epididymis. While 27 the change in tsRNA-Leu-CAA was temporary and did not persist, strikingly tsRNA-Arg-CCT-28 2 deregulation persisted until 14 days post injection (Figure 1C). To additionally validate the

1 Dex induced change of tsRNA-Arg-CCT-2 independent of epididymal transit from caput to 2 cauda we replicated the effect observed in mature sperm sampled from cauda epididymis 3 3 hours post injection using q-PCR (Figure 2D, Supplementary table 2). Additionally, we sampled 4 caput sperm 3 hours post injection and measured tsRNA-Arg-CCT-2 levels. An overall 2-way 5 ANOVA revealed a significant interaction between sperm sampling location (caput versus 6 cauda) and treatment (vehicle versus Dex). Post hoc tests confirm a significant increase in 7 tsRNA-Arg-CCT-2 levels in response to Dex in cauda but not in caput sperm and a significant 8 increase in Arg-CCT-2 levels between cauda and caput sperm independent of treatment.

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10 The behaviour of miRNAs differed considerably from tsRNAs. As would be expected if 11 epididymal transit was required for miRNA changes to be implemented, we observe no group 12 effect of treatment on miRNAs (Supplementary Fig. 2A, Supplementary table 2) across 3 hours 13 and 7 days post injection. Further, we detected no effect of time post injection on sperm miRNA 14 payload (Supplementary Fig. 2B, Supplementary table 2) confirming the absence of an effect 15 of injection on miRNAs per se. However, we neither detected an interaction between Dex and 16 time post injection (Supplementary Fig. 2C, Supplementary table 2) in miRNAs 7 days and 3 17 hours post injection. Importantly, when inspecting those miRNAs that were significantly altered 18 14 days after injection, no alterations were apparent 3 hours or 7 days post injection 19 (Supplementary Fig. 2D), indicating that changes in miRNAs occur more slowly or require 20 sperm cells to reside in the testes at the time of injection.

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22 Effects of acute GR activation on in vivo offspring metabolic phenotype

Based on the two reports on effects of single foot shock on offspring weight and the impact of a single GR activation on germline small RNA payload, we hypothesized that this acute impact on the receptor is sufficient to elicit intergenerational effects. We thus injected Dex once intraperitoneally, then harvested sperm 14 days post injection, and performed *IVF* using naïve oocytes to generate offspring for phenotyping (Figure 3A). Dex treatment did not affect sperm count, fertility rate or resulting litter-sizes (Supplementary Fig.6).

1 The weight and size of pups was measured every 2 to 4 weeks starting at weaning (3 weeks 2 of age) until adulthood (12 weeks of age) and body mass index (BMI) was calculated as a ratio 3 of weight and squared length. Overall ANOVA of the resulting offspring showed a significant 4 effect of treatment ((F1.71)= 76.55, p<0.0001), time post injection ((F2.087,144.7)=41.99, 5 p<0.0001) and sex (F (1, 71) = 76.55, p<0.0001) on BMI, and a significant interaction between time and sex (F (3, 208) = 33.75, p<0.0001) and time and treatment (F (3, 208) = 5.834, 6 7 p=0.0008) (Figure 3B, Supplementary Fig. 3C, Supplementary table 3). These results show 8 that while males had generally higher BMI, both male and female offspring of Dex injected 9 fathers had a higher BMI.

10 To further explore potential causes and consequences of altered BMI, adult animals were 11 additionally tested for their glucose tolerance following glucose injection. Overall ANOVA 12 analysis of blood glucose levels revealed a significant effect of sex (F (1, 44) = 54.80, 13 p<0.0001) and time post injection (F (2.593, 114.1) = 196.6, p<0.0001) and significant 14 interactions between sex and time post injection (F (4, 176) = 6.115, P=0.0001), and sex and treatment (F (1, 44) = 15.62, P=0.0003) (Supplementary Fig. 3C). Follow up repeated 15 16 measurements ANOVA analysis separated by sex showed a significant effect of treatment. 17 time and interaction in females (treatment: F (1, 22) = 12.35, p=0.0020; time: F (4, 88) = 110.1, 18 p<0.0001; interaction: F (4, 88) = 2.835, p=0.0291) and significant effects of treatment and time but no interaction in males (treatment: F (1, 22) = 6.019, p=0.0225; time: F (4, 88) = 96.36, 19 20 p<0.0001; F (4, 88) = 0.5401, p=0.7067; Figure 3C). These data hence demonstrate a sex-21 dependent effect of paternal Dex injection on glucose tolerance, with impaired tolerance in 22 females and decreased glucose levels in males in response to glucose challenge.

In addition, blood glucose levels were assessed during the insulin tolerance test. Overall
 ANOVA analysis showed significant effects of sex (F (1, 37) = 162.6, P<0.0001)

and time (F (3.314, 122.6) = 23.85, P<0.0001) and revealed a significant interaction between
sex and time (F (4, 148) = 12.49, P<0.0001), time and treatment (F (4, 148) = 5.380, P=0.0005)
and time and treatment and sex (F (4, 148) = 5.392, P=0.0004) (Supplementary Fig. 3C).
Follow-up repeated measurements ANOVA separated by sex showed a significant effect of

1 time (F (2.982, 65.60) = 44.73, p<0.0001), yet no significant effect of treatment (F (1, 22) = 2 0.3465, p=0.5621) nor an interaction between time and treatment (F (4, 88) = 0.1373, 3 p=0.9681) in females (Figure 3D). In males we observe no effect of treatment (F(1, 15) = 1.467, 4 p=0.2446) yet detected a significant effect of time post injection time (F(2.914, 43.71) = 4.538, 5 p=0.0079), and a significant interaction between treatment and time post injection (F(4, 60) = 6 7.003, p=0.0001, Figure 3D). These results indicate sex and time dependent effects of 7 paternal Dex on insulin tolerance. They further show no change in insulin tolerance in female 8 descendants of fathers injected with Dex, yet impaired insulin tolerance in male progeny.

9 Lastly, we explored a potential reflection of altered BMI in tissue composition by necropsy and 10 weighing the dissected organs and fat pads. Overall ANOVA of necropsy weights revealed a 11 significant effect of sex (F (1, 140) = 28.27, P<0.0001), tissue (F (4, 140) = 232.7, P<0.0001) 12 and a significant interaction between sex and tissue (F (4, 140) = 3.379, P=0.0113) yet no 13 effect of treatment (F (1, 140) = 0.2587, P=0.6118), or interaction between treatment and sex 14 (F (1, 140) = 0.0004794, P=0.9826) or treatment and tissue (F (4, 140) = 0.1635, P=0.9565) 15 on tissue weight (Supplementary Fig. 3A,B,C). This confirms sex dependency, yet no effect of 16 paternal Dex injection on tissue weight in both sexes.

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18 Effects of acute Dex on offspring early embryonic small RNA

19 The small quantity of paternal RNAs in the zygote relative to the large pool of maternal RNAs 20 poses serious obstacles to their accurate quantification (Chen, Yan and Duan, 2016). While 21 initial reports on small RNA transmission relied on comparative sequencing or microarray 22 analyses of unfertilized oocytes and fertilized zygotes (Ostermeier et al., 2004), today we are 23 aware that such comparisons can be deceiving, as they rely heavily on both assessment 24 method (e.g. microarray restricted to a selective set versus unbiased genome-wide 25 sequencing) and sequencing depth (Dard-Dascot et al., 2018; Yeri et al., 2018). One such 26 example is inconsistent results regarding miRNAs that are exclusively supplied from the 27 sperm, such as miR-34c, -99a, -214 (Amanai, Brahmajosyula and Perry, 2006; Liu et al., 2012). 28 Alternative approaches have used indirect measures, e.g. assessing mRNA targets of

1 paternally derived small RNAs (Amanai, Brahmajosyula and Perry, 2006; Tang et al., 2007; 2 Krawetz et al., 2011; Sharma et al., 2015). We attempted to directly examine the relative 3 difference between the small RNA landscape in early embryos resulting from IVF of naïve 4 oocytes with sperm from either Dex or vehicle injected males (Figure 4A). We used small-RNA 5 sequencing to compare 2-cell embryos derived from Dex treated or control fathers. We 6 detected an average of 29 % mappable reads including 21% multimappers. While we only 7 detected subtle changes in miRNAs of Dex exposed progeny, we observed downregulation of 8 several tsRNAs from 6 different genomic locations (q<0.1) (Figure 4B). Strikingly, two of the 9 downregulated tsRNAs (Gly-GCC at several genomic loci and Gly-CCC) were consistently 10 downregulated in sperm 14 days post Dex injection. This could either indicate a reduced 11 delivery of this sperm RNA cargo in Dex treated males to the oocytes they fertilize, or earlier 12 usage and function of respective RNA in Dex leading to a quicker elimination or shorter half-13 life.

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16 Effects of acute Dex administration on offspring early embryonic transcriptome

17 If sperm RNA were directly impacting the zygotic mRNA pool or if they were affecting early 18 embryonic gene expression, then this should be apparent in the 2 cell embryo's transcriptome 19 (Figure 5A). To examine the effect of paternal Dex on early embryonic RNA content we 20 subjected 2-cell embryos to the Smartseg single cell sequencing protocol (Supplementary Fig. 21 4). After performing quality control and filtering the sequenced 2-cell embryo data on criteria 22 such as minimal read count/embryo (Supplementary Fig. 4A), we carried out unsupervised 23 clustering based on their gene expression profiles using SC3 (Kiselev et al., 2017). We 24 identified two distinct clusters (C1 and C2), which were composed by a balanced mixture of 25 treated and control cells. (Figure 5B). Since the resolution of single cell experiments allows 26 characterizing distinctive transcriptomic profiles within early cell division stages, we used 27 scmap (Kiselev, Yiu and Hemberg, 2018), to project each 2 cell embryo gene expression 28 profile onto a reference dataset of single cells from 2 cell embryo states previously reported

by Deng *et al.* (Deng *et al.*, 2014) (Figure 5C.) Most of the 2 cell embryos belonging to cluster
 C1 projected to the late 2 cell stage, whereas embryos from C2 exclusively projected to cells
 from the mid 2 cell stage. This shows that the two clusters identified through unsupervised
 clustering correspond to 2-cell embryos in the mid and late 2-cell stage respectively.

5 Principal component analysis (PCA) revealed a prominent separation between C1 and C2 6 along the PC1 axis, suggesting a correlation between PC1 and developmental transitions 7 between mid and late single cell embryos (Supplementary Fig. 4B.) Interestingly, 2 cell embryo 8 offspring of males injected with Dex exhibited a significant shift of the C1 cluster across PC1 9 (two-sided Wilcox test p< 0.03), while the C2 clusters did not show significant differences 10 across PC1 between treatment and control groups (Figure 5D.) These results suggest that the 11 effect of paternal Dex treatment on the transcriptome only becomes apparent at the late 2 cell 12 embryo stage. To further explore this hypothesis, we calculated the silhouette coefficient 13 (Rousseeuw, 1987) on PC1, as a measure of distance between C1 and C2 clusters, for the 14 control and treatment group. We observed a significant increase of PC1 silhouette coefficient 15 between treatment and controls for both C1 (one-sided Wilcoxon test p-value <0.005) and C2 16 (one-sided Wilcoxon test p-value $< 2 \times 10^{-5}$.) This confirms that Dex treatment affects 17 embryonic gene expression, promoting altered late 2 cell embryo stages since the divergence 18 from mid 2 cell embryos is significantly bigger in Dex offspring compared to control offspring 19 (Figure 5E.)

20 Accordingly, differential gene expression analysis using Monocle2 (Qiu et al., 2017) focused 21 on late 2-cell embryos (cluster C1) revealed significant gene expression changes between 22 offspring of males injected with Dex and controls across 38 genes, some of which were already 23 apparent to a less significant extent during mid-2 cell embryos (cluster C1; e.g. Tcl1; 24 Supplementary Table 4 In line with a potentially altered developmental trajectory becoming 25 apparent in cluster 1, the late 2-cell stage includes several affected genes that are involved in 26 early embryonic development. For example, Bcap31 (B-cell receptor-associated protein 31) is 27 an important element for endoplasmic reticulum and Golgi apparatus function, and Bcap31 28 mutations lead to developmental diseases with metabolic disturbances (Cacciagli et al., 2013).

1 This is reminiscent of the metabolic phenotype observed in the adult offspring of Dex injected 2 fathers. Hprt (Hypoxanthine-guanine phosphoribosyltransferase) is crucial for cell cycle 3 division, and Tcl1 (T-cell leukemia/lymphoma) regulates cell proliferation (Kang et al., 2013; 4 Miyazaki et al., 2013). Hence, an upregulation of Hprt and a concomitant down-regulation of 5 Tcl1 might indicate that cell fate decisions later during development may be affected. Another 6 differentially expressed gene is Rbbp7 (RB binding protein 7), which is part of many histone 7 deacetylase complexes such as Nurd and PRC2/EED-EZH2, and thus plays an essential role 8 in chromatin mediated gene regulation (Yu et al., 2018). Interestingly, several forms of PRC 9 mutations in humans lead to different kinds of overgrowth phenotypes (Deevy and Bracken, 10 2019), an abnormality reminiscent of the increased BMI observed in Dex-offspring (Figure 3B.)

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13 Effects of Dex administration on an interesting candidate for sperm RNA mediated inheritance 14 Despite the observed changes in sperm tsRNAs following acute Dex injection, we did not find 15 an obvious causal connection to the altered 2-cell embryonic transcripts. This prompted us to 16 investigate whether other germline changes might be more crucial for the offspring's in vivo 17 alterations in our model. We previously showed that chronic stress exposure also led to 18 changes in sperm long RNAs that contributed functionally to the transmission of effects to the 19 offspring (Gapp et al., 2018). Yet the fact that sperm RNA is stable through transmission and 20 that the minute amounts of transmitted paternal RNA can elicit major changes in the embryo 21 remains puzzling. Therefore, we evaluated the impact of Dex injection on the highly stable 22 class of circRNAs in male sperm. CircRNAs were previously detected in swine(Godia et al., 23 2020) and human sperm (Chioccarelli et al., 2019) and suggested to have functional 24 implications in epigenetic regulation. They have been attributed a critical role in the male 25 germline after cessation of transcriptional activity (Tang et al., 2020). Using Circexplorer in 26 combination with EdgeR, analysis of sperm long RNA sequencing of males treated with Dex 27 and controls revealed significant upregulation of two circRNAs (Figure 6A, q<0.1) and we also 28 observed several significant changes in the sperm long RNA protein coding transcripts

1 following acute Dex treatment (Figure 6B,C, Supplementary Fig. 1B, Supplementary table 5). 2 Both circRNAs are hosted in genes relevant for immune function (Taspase 1: Tasp1 and DENN 3 Domain Containing 1B Dennd1b), yet the host genes did not show differential abundance 4 of the protein coding transcript (Supplementary table 5). We then replicated the upregulation 5 of these to CircRNAs by q-PCR in cauda epididymal sperm of a separate set of animals using 6 CircRNA specific primers that span the backsplice junction. At the same time, we also 7 assessed their abundance in caput epididymal sperm to evaluate whether the observed effect 8 was also present in sperm cells under development. We observed a significant interaction 9 between treatment and tissue. Post hoc tests confirm the increased abundance of CircTasp1 10 and CircDend1 in cauda sperm detected in the sequencing analysis. No change was 11 detectable in caput sperm, arguing against an induction of the change in developing sperm 12 cells (Figure 6D,E). Given the absence of transcription, these data suggest that in control 13 conditions these CircRNAs exert their function during spermiogenesis (for example by being 14 translated) and that by contrast in Dex treated cells the CircRNA is not consumed to the same 15 extend leading to an apparent upregulation in mature cauda epididymal sperm. CircAtlas(Wu, 16 Ji and Zhao, 2020) revealed several potential miRNA sponge-targets to be captured by the 17 altered circRNAs. Some of these miRNAs are common sponge-targets of both circRNAs such 18 as mir3110-5p, mir706, mir1955 (Supplementary Fig. 5). Diamine acetyltransferase 1 (Sat1), 19 one of 3110-5p's high confidence miRNA-targets, as predicted by TargetScan(Agarwal et al., 20 2015), is indeed significantly upregulated in the embryos composing cluster 1 (later 21 developmental stage). MiRNA-target-upregulation is expected if mir3110-5p was 22 downregulated through circRNA-mediated-sponging and highlights a potential effective 23 contribution of increased circRNA in sperm to embryonic pathway regulation. This is the first 24 report of a change induced by environmental exposure in this compelling class of RNA in 25 sperm.

- 26
- 27 Discussion

1 By generating offspring using assisted reproductive techniques (IVF), we circumvent potential 2 confounding variables such as transmission via RNA contained in seminal exosomes (Vojtech 3 et al., 2014) and affected maternal care by altered mating behavior (Mashoodh et al., 2012) 4 and prove germline dependence (Bohacek and Mansuy, 2017). Consistent with the significant 5 changes of miRNAs and tsRNAs in the germline 2 weeks post GR activation, previous studies 6 including our own have observed regulation of mouse sperm small RNA in a variety of contexts 7 (Rodgers et al., 2013; Gapp et al., 2014; Grandjean et al., 2015; Sharma et al., 2015; Chen et 8 al., 2016; Wu et al., 2016; Short et al., 2017; Benito et al., 2018; Rompala et al., 2018; 9 Yeshurun and Hannan, 2018).

10 Especially relevant specifically for our analysis, sperm RNA sequencing after drinking water 11 administration of corticosterone for 4 weeks followed by mating, led to strong downregulation 12 of tsRNA-GluCTC and tsRNA CysGCA, two of our top down-regulated tsRNAs, indicating that 13 these tsRNAs are responding similarly to acute and chronic insults. At the same time this 14 chronic manipulation elicits changes of several miRNAs, e.g. 34c and 471 (Short et al., 2016), 15 albeit in the opposite direction of what we find in response to acute Dex treatment. These 16 discrepancies may arise either by the Dex induced short-term suppression of internal 17 corticosteroid (Barden, 1999) or due to adaptations in response to chronic administration.

While four (Petropoulos, Matthews and Szyf, 2014; Short *et al.*, 2016; Wu *et al.*, 2016; Cartier *et al.*, 2018) out of five (Bönisch *et al.*, 2016) previous studies did report phenotypic effects following chronic paternal Dex exposure, only two assessed sperm small RNAs to associate the alterations to the sperm RNA payload (Bönisch *et al.*, 2016; Cartier *et al.*, 2018) yielding conflicting outcomes.

These differences might be due to inconsistent life stages (adulthood versus gestational), sperm collection (swim up, somatic lysis, or no purification) and/or dosage of exposure. Depending on the dosage and timing, the complex autoregulation of the GR can lead to GR downregulation after prolonged activation (Gjerstad, Lightman and Spiga, 2018). Acute exposures have the advantage of avoiding such long-term feedback regulation, and hence provide an elegant approach for studying the signaling pathways leading to germline changes.

1 Mature sperm tsRNAs and miRNAs have been shown to be acquired during epididymal transit 2 (Sharma et al., 2015, 2018) and miRNAs are necessary for early embryonic development 3 under certain circumstances (Liu et al., 2012; Conine et al., 2018)(Zhou et al., 2019)(Wang et 4 al., 2020). Furthermore, a recent publication suggests that chronic stress induced sperm 5 miRNAs are taken up primarily from epididymosomes originating from the caput epididymis or 6 the proximal epididymal tract (Chan et al., 2020). Chronic nutritional manipulation with effects 7 on offspring also suggest the necessity of epididymal transit to acquire tsRNA changes in 8 sperm(Sharma et al., 2015). Harvesting mature sperm 3 hours after exposure yields a 9 population enriched for cells that had been exposed while already in the cauda epididymis, 10 where spermatids reside for roughly 5 days (Meistrich, 1975). These cells have not traveled 11 through the epididymis nor have they had a chance to potentially take up small RNA from 12 caput-derived epididymosomes after Dex administration. As expected, we detect no changes 13 in miRNAs in these samples. We do however detect changes in tsRNAs 3 hours post Dex, 14 some of which even persist 14 days post injection. These results show rapid acquisition of 15 changes in vivo and corroborate previous in vitro findings that show that incubation with 16 epididymosomes can alter sperm RNA payload(Sharma et al., 2015). Our acute intervention 17 assesses effects on germline payload already after a short interval, whereas chronic 18 interventions - based on their experimental design - do not assess changes in mature sperm 19 soon after the first intervention. Studies aiming at the elucidation of the origin of sperm RNA 20 changes might benefit from acute interventions to circumvent confounders such as dynamic 21 exosomal RNA supply as a result of cumulative interventive strain on animals.

An additional option for sperm RNA alterations in transcriptionally inert sperms was suggested in a recent study that found mitochondrial tRNA cleavage in the T-loop in response to a oneweek high sugar diet in humans(Nätt *et al.*, 2019). In line with this observation, Dex injection could trigger oxidative stress(Bera *et al.*, 2010) which provokes such cleavage to increase tsRNA levels(Thompson *et al.*, 2008). A role for oxidative stress in sperm RNA dynamics is further supported by a recent study in boar sperm that found seasonal differences in sperm small and long RNA associated with changed abundance of transcripts mapping to oxidative

stress-, DNA damage- and autophagy- related genes (Gòdia *et al.*, 2019). However, such
 potentially oxidative stress mediated mechanism does not explain a rapid decrease of tsRNAs
 3 hours post Dex injection.

4 Importantly, we show alterations in sperm tsRNAs that persist in the oocyte concomitant with 5 changes in early embryonic gene expression and a metabolic phenotype in adulthood. tsRNAs 6 and tRNA-Gly derived fragments in particular are known to induce chromatin structure 7 mediated gene regulation and to regulate cell differentiation in various contexts (Li et al., 2016; 8 Guzzi and Bellodi, 2020). Hence, we propose that the transmitted reduction in key tsRNAs 9 such as Gly-GCC and Gly-CCC explains in part the observed perturbations during late two cell 10 embryo developmental stage. This might reflect an accelerated developmental transcriptional 11 program in the preimplantation embryo of Dex injected males, ultimately resulting in aberrant 12 BMI and glucose metabolism later in life.

13 Additionally, we have discovered alterations in circRNA abundance in mature sperm that might 14 also impact the developmental program in the early embryo. CircRNAs have the potential to 15 be translated into proteins via back-splicing(Legnini et al., 2017). Accordingly, they are crucial 16 contributors to spermiogenesis post transcriptional cessation, since they provide a stable 17 alternative to linear mRNA templates for protein translation(Tang et al., 2020). Transmitted 18 sperm-circRNAs could likewise contribute to translation post fertilization, yet the 19 unconventional lattice-state of ribosomes preventing normal rates of translation post 20 transcription(Israel et al., 2019) accompanied by a rapid increase in proteins of the 21 ubiquitine/proteasome pathway(Wang et al., 2010) make this unlikely. Nevertheless, a study 22 on human sperm detected abundant levels of circRNA with predicted regulatory function of 23 early developmental genes in sperm heads, suggesting transmission and function post 24 fertilization(Chioccarelli et al., 2019). By sponging miRNAs that regulate early embryonic 25 transcripts, circRNA could amplify minute signals from paternal environment, such as might be 26 the case for the gene transcript Sat1, that displays increased expression in Dex offspring in 2-27 cell embryos from cluster 1.

Besides altered RNA identity, nucleic acid modifications especially of RNA but also DNA methylation and chromatin accessibility might contribute further to the effects of Dex injections on offspring metabolism. While detection of changes in each player should be subject of further investigation and might reveal a glimpse of their potential implication, proof of the individual relative causal contribution is extremely challenging since they likely require tight interaction to unravel their orchestrated effects.

7 Finally, it might be useful to consider testing the translatability of our findings to humans. Here 8 we investigated the effects of a single Dex administration soon after the injection in mice, 9 mimicking a single GR activation such as elicited by treatment of an acute asthma 10 exacerbation (Cross, Paul and Goldman, 2011). The recent report that Dex can reduce the 11 number of deaths associated with the Covid-19 pandemic ('Biggest COVID-19 trial tests 12 repurposed drugs first', 2020), further prompts the re-evaluation of the impact of prolonged 13 Dex treatment on offspring phenotype. From a clinical perspective, additional consideration is 14 warranted for consequences on offspring health when extended time has elapsed between 15 treatment and time of conception. Such designs may pave the way for the extrapolation of our 16 findings.

17 We conclude that acute Dex treatment can induce germline epimodifications in the form of 18 small and long non-coding RNA, which likely are relevant in the transmission of the effects of 19 single traumatic events on offspring well-being. Our data suggest that sperm small RNAs are 20 not solely regulated via epididymosomal uptake during transition from caput to cauda 21 epididymis. This expands the interpretation from chronic dietary and stress exposures(Sharma 22 et al., 2015; Chan et al., 2020), where uptake of tsRNAs and miRNAs via epididymosomes 23 has been suggested to lead to differential sperm payload, yet required sperm to transit from 24 caput to cauda to bring about the changes. A persistent detection of significant fold changes 25 of the exact same sperm small RNA in the embryo suggests functional implication in the 26 information transfer from father to offspring. Together with potentially transmitted miRNA 27 sponges in the form of circRNAs, this likely contributes to a slight developmental acceleration 28 of gene expression programs in the early embryo and ultimately manifests in a metabolic

phenotype. Future studies may aim at testing the causal contribution of specific sperm RNAs to the transmission of effects of acute impacts. Certainly, continuous methodological refinement will help dissect the relative implication and the interplay of the distinct germline modifications such as DNA-methylation, histone-PTMs and chromatin architecture in this highly complex process.

6

7 Limitations of study

8 This study identifies a highly dynamic response of sperm RNA cargo in response to a single 9 Dex administration and presents altered CircRNAs in mature sperm in response to an 10 environmental insult with consequences on the progeny's metabolism. While our study implies 11 a functional role of altered sperm RNAs in the transmission of a Dex induced phenotype to the 12 offspring a causal proof will require RNA injections into fertilized naïve oocytes. Second, 13 although we identified one tsRNA to be persistently altered in serum exosomes as well as in 14 caput and cauda sperm and at different times post Dex injection, the identification of the tissue 15 of origin of the altered tsRNAs would require technically highly challenging metabolic labelling 16 experiments.

17

18 Author's contributions

KG performed animal exposures, collected samples and prepared sequencing libraries. JK helped with sample collection and processing and performed q-PCRs. EG performed IVF and embryo culture. GP analysed single cell sequencing data. MH supervised the single-embryo sequencing analysis. AC and FG analyzed bulk sequencing data. KG, JB and EAM designed the study, interpreted the results and wrote the manuscript with input from the other authors.

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25

26 Acknowledgements

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14 **Declaration of interests**

15 The authors declare no competing interests. E.A.M. is a founder and Director of STORM 16 Therapeutics Ltd. STORM Therapeutics had no role in the design of the study and collection, 17 analysis, and interpretation of data as well as in writing the manuscript.

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1 Figure legends

2 Figure 1 Effects of Dex on small RNA payload of sperm cells residing in testis at time of 3 administration. (A) Experimental design depicting a time window of two weeks between 4 injection of Dex and sperm collection for molecular analysis. (B) Volcano plot depicting fold 5 changes and significance level of miRNAs and tsRNAs (C) in mature sperm 14 days post 6 injection of Dex (n=8) versus vehicle (n=8) as assessed by small RNA sequencing. (D) Stacked 7 barplot showing reads of different sizes mapping to rRNAs in Dex (n=8) and vehicle (n=8). (E) 8 Volcano plot demonstrating global increased abundance of rRNA fragments. (F) Q-PCR 9 results of small RNA assays for caput sperm (tsArg-CCT-2: t(49)=3.49, g<0.001, tsGlyGCC: 10 t(49)=0,32, q>0.05; tsHIs-GTG: t(49)=0.29, q>0.05; tsLeu-CAA-2: t(49)=0.62, q<0.5; mir34c-11 5p: t(49)=0.27, q>0.05; mir6538: t(49)=0.59, q>0.05, mir677-5p t(49)=0.62 q>0.05) (G) serum 12 exosomes (tsArg-CCT-2: t(33)=3.43, q<0.01, tsGlyGCC: t(33)=0,04, q>0.05; tsHIs-GTG: 13 t(33)=0.07, g>0.05; mir34c-5p: t(33)=0.51, g>0.05; mir6538: t(33)=0.02, g>0.05) days after 14 drug injection. Whiskers display minimum and maximum. **q<0.01, ***q<0.001 multiple t-tests 15 corrected for multiple testing.

16

17 Figure 2 Effect of Dex on sperm cells at different time points post Dex administration. (A) 18 Experimental design showing the location of sperm at time of injection and timing of sperm 19 harvest. MA (log-intensity ratios (M-values) versus log-intensity averages (A-values) plots 20 depicting (B) effect of Dex (log2 fold changes control versus dexamethasone), (C) of time post 21 injection (log2 fold changes 7 days versus 3 hours) (7 days Dex n =4 and controls n =4, 3 22 hours Dex n = 3 and controls n = 4). TsRNAs are indicated by sequence identity for display only, 23 each dot represents one small RNA. MA plot depicts log2 fold changes on the y axis and the 24 expression level on the x axis (the higher the expression the further to the right). Statistically 25 significantly changed small RNAs are highlighted in red q<0.05. (D) Relative expression of 26 ArgCCT-2 as obtained by q-RT-PCR (cauda: Dex n=4, controls n=4, caput: Dex n=4, controls 27 n=5; interaction F(1,13)=6.34, p=0.0257, treatment F(1,13)=5.97, p=0.0040, site of collection 28 (F(1,13)=12.15, p=0.0296); cauda control versus cauda Dex t(13)=3.42, p=0.0274, cauda Dex

verus caput Dex t(13)=4.137, p=0.007). Whiskers display minimum and maximum. *p<0.05,
 mixed effect model group effect of location (REML), **p<0.01 multiple comparisons SIDAK
 corrected.

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6 Figure 3 Effect of Dex on metabolic phenotype in the offspring (A) Experimental design 7 depicting timeline between injection, sperm harvest, in vitro fertilization and phenotyping. (B) 8 Impact of Dex on male and female adult offspring (B) Body mass index (males vehicle offspring 9 n=21, Dex offspring n=22, females vehicle offspring n=17, Dex offspring n=17) (C) glucose 10 tolerance (males vehicle offspring n=12, Dex offspring n=12, females vehicle offspring n=12, 11 Dex offspring n=12) and (D) insulin tolerance (males vehicle offspring n=9, Dex offspring n=8, 12 females vehicle offspring n=12, Dex offspring n=12). Error bars display SEM. Detailed 13 statistical results are depicted in Supplementary Fig. 5, raw data are provided in supplementary table 3). *p<0.05, ** p<0.01, ***p<0.001, 3-way repeated measures ANOVA. 14

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Figure 4 Effect of paternal Dex injection on embryonic offspring small RNA (A) Experimental design depicting timeline between injection, sperm harvest, in vitro fertilization and small RNA sequencing at 2-cell stage. (B) Heatmap showing effect of paternal Dex on small RNA tsRNAs (vehicle embryonic offspring n=5, Dex embryonic offspring n=4). TsRNAs are grouped by sequence identity for display only.

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Figure 5 Effect of paternal Dex injection on embryonic offspring long RNA transcriptome (A) Experimental design depicting timeline between injection, sperm harvest, in vitro fertilization and Smartseq2 sequencing at 2-cell stage. (B) Consensus matrix representing the similarity between cells as reported by SC3. Similarity 0 indicates that a given pair of embryos were never assigned to the same cluster, whereas similarity 1 means that a pair of embryos were always assigned to the same cluster. (C) Sankey diagram showing projection of the obtained

1 clusters (C1 and C2) into clusters reported by Deng et al. for single cells obtained from 2 cell 2 embryos. (D) Principal component analysis of two-cell embryos gene expression. The top 3 panel indicates the density of 2 cell embryos along PC1 grouped by condition; control (red) 4 and Dex treatment (blue). The two bottom panels show the distribution of 2 cell embryos across 5 PC1 and PC2 for control (red) and treated (blue) groups. The cluster membership of each 6 embryo is denoted by the point shapes (C1 cycles; C2 triangles) and the centroids of each 7 cluster is indicated with a black symbol overlaid with an x. Wilcox tests were performed to 8 assess differences on PC1 values of C1 and C2 clusters between the treated and control 9 groups. NS denotes non-significant change for C2 cluster, while * indicates a significant 10 difference for C1 cluster (p-value< 0.05). (E) Silhouette coefficient comparison between 11 treatment and control, statistical significance was assessed with Wilcox test (** p-value < 0.01; 12 *** p-value < 0.005) (F) Selection of differentially expressed genes as determined by Monocle 13 within C1 corresponding to late 2-cell embryo stage (*** adjusted p-value < 0.005).

Figure 6 Effects of Dex on long RNA payload of sperm cells residing in testis at time of administration. (A) Experimental design depicting a time window of two weeks between injection of Dex and sperm collection for molecular analysis.

17 (B) Volcano plot depicting fold changes and significance level of long RNA in mature sperm 18 14 days post injection of Dex (n=4) versus vehicle (n=4) as assessed by small RNA 19 sequencing. Statistically significant transcripts are highlighted in red (FDR < 0.05). (C) 20 Heatmap showing significantly differentially expressed long RNA transcripts of the same 21 experiment (multiple comparison corrected, q<0.05). (D) Q-PCR results of CircRNA assays for 22 caput and cauda sperm for Tasp1 (interaction treatment x tissue F(1,5)=7.53, p<0.05, caput: 23 control n=4, DEX n=5, p>0.05; cauda control n=5, DEX n= 3, p<0.01) (*p<0.05 mixed effect 24 model interaction (REML), **q<0.01 multiple comparisons SIDAK corrected) and Dennd1 25 (interaction treatment x tissue F(1,15)=6, p<0.05, caput: control n=4, DEX n=5, p>0.05; cauda 26 control n=5, DEX n= 5, p<0.05) (E). Whiskers display minimum and maximum. *p<0.05 mixed 27 effect model interaction (REML), *q<0.05 multiple comparisons SIDAK corrected.

Timepoint post injection	3 hours	7 days		14 days	
miRNAs	Cauda	Cauda	Cauda	Caput	Serum
3535	no ch.	no ch.		not det	not det
677-5p	no ch.	no ch.	1	no ch.	not det
677-3p	no ch.	no ch.	1		
34c-5p	no ch.	no ch.	\downarrow	no ch.	no ch.
7033-5p	no ch.	no ch.	1		
703	no ch.	no ch.	1		
5126	no ch.	no ch.	1		
5121	no ch.	no ch.	1		
6240	no ch.	no ch.	1		
5114	no ch.	no ch.	1		
1839-3p	no ch.	no ch.	1		
1949	no ch.	no ch.	1		
196a-5p	no ch.	no ch.	↑		
3064-5p	no ch.	no ch.	1		
196b-5p	no ch.	no ch.	↑		
6538	no ch.	no ch.	1	no ch.	no ch.
18a-5p	no ch.	no ch.	\downarrow		
3963	no ch.	no ch.	\downarrow		
471-5p	no ch.	no ch.	\downarrow		
5099	no ch.	no ch.	1		
1843b-3p	no ch.	no ch.	1		
1843a-3p	no ch.	no ch.	1		
circRNAs					
Tasp1			1	no ch.	
Dennd1b			1	no ch.	

1 Table 1 Changes in miRNAs and CircRNAs across time and sample type

Table 1: Table depicts significantly altered small RNAs and circRNAs in cauda sperm at 14days post dexamethasone injection.

Table 2 Changes in tsRNAs across time and sample type

Timepoint post injection	3 hours		7 days		14 days		
tsRNAs	Cauda	Caput	Cauda	Cauda	Caput	Serum	
Leu-CAA	1		1	no ch.	no ch		
Thr-TGT1	Ļ		1	no ch.			
His-GTG-2	no ch		Ļ	no ch	no ch	no ch.	
His-GTG-3	no ch		↑	no ch.			
Pro-AGG-1	no ch		1	no ch.			
Pro-TGG-2	no ch		↑	no ch.			
Pro-TGG-4	no ch		↑ ↑	no ch.			
Glu-CTC-4	no ch		1 ↑	no ch.			
Pro-CGG-1	no ch		 ↑	no ch.			
Gly-GCC-6	no ch		1	no ch			
Gly-GCC-2	no ch		no ch		no ch	no ch	
Arg-CCT-2		no ob		↓		-	
•	1	no ch.	T	T	1	1	
Ala-TGC-2	no ch		no ch	\downarrow			
Arg-ACG-3	no ch		no ch	\downarrow			
Glu-CTC-3	no ch		no ch	\downarrow			
Ser-TGA-1	no ch		no ch	\downarrow			
Thr-TGT-2	no ch		no ch	\uparrow			
Ser-TGA-2	no ch		no ch	1			
Ser-AGA-1	no ch		no ch	1			
Glu-TTC-3	no ch		no ch	\downarrow			
Arg-CCT-1	no ch		no ch	1			
Ser-AGA-2	no ch		no ch	1			
Cys-GCA-3	no ch		no ch	1			
Pro-AGG-3	no ch		no ch	↑			
Lys-CTT-3	no ch		no ch	1			
Arg-CCT-3	no ch		no ch	¥			
Lys-CTT-3	no ch		no ch				
Trp-CCA-5	no ch		no ch	+			
Cys-GCA-2	no ch		no ch				
Ala-TGC-5	no ch		no ch	¥			
Gly-CCC-3	no ch		no ch	+			
Gly-CCC-4	no ch		no ch	¥			
Ala-TGC-5	no ch		no ch	¥			
Thr-AGT-5	no ch		no ch	↓ 			
SeC-TCA-1	no ch		no ch				
Thr-AGT-7	no ch	-	no ch	↓		-	
Glu-CTC-2							
His-GTG-1	no ch no ch		no ch no ch	↓			
Arg-CCT-4	no ch		no ch				
Lys-CTT-1	no ch		no ch	↓	+	+	
Lys-CTT-2	no ch		no ch	↓			
Lys-CTT-2	no ch		no ch	<u>↓</u>			
Asp-GTC-4	no ch		no ch	↓			
Gly-GCC-5	no ch		no ch	↓ ↓			
Thr-TGT-3	no ch		no ch	1			
Asn-GTT-2	no ch		no ch	1			
Asn-GTT-4	no ch		no ch	↑			
Asn-GTT-1	no ch		no ch	1			
Gly-GCC-4	no ch		no ch	\downarrow			
Gly-GCC-2	no ch		no ch	\downarrow			
Thr-CGT-4	no ch		no ch	1			
Gly-CCC-5	no ch		no ch	Ļ			
Ala-TGC-6	no ch		no ch	, i i i i i i i i i i i i i i i i i i i	1	1	
Glu-CTC-1	no ch	1	no ch	Ť	1	1	

- **Table 2:** No highlight: Significantly altered in cauda sperm at 14 days post dexamethasoneinjection. In bold: Persistently altered in cauda sperm at 7 days and 3 hours. Italic: Significantinteraction between 7 days and 3 hours.
- 2 3 4

1 STAR METHODS

- 2 Detailed methods are provided in the online version of this paper and include the following:
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1	STAR METHODS					
2	RESOURCE AVAILABILITY					
3	Lead contact					
4	Further information and requests for resources and reagents should be directed to, and will be					
5	fulfilled upon reasonable request, by the Lead Contact, Katharina Gapp					
6	(katharina.gapp@hest.ethz.ch).					
7						
8	Materials availability					
9	The study did not generate new reagents.					
10						
11	Data and code availability					
12	- The raw datasets supporting the conclusions of this article are included within the					
13	article (supplementary tables).					
14	- Sequencing data have been deposited at Gene omnibus and ENA and are publicly available					
15	as of the date of publication. All sequencing data were deposited to Gene Omnibus and					
16	ENA.					
17	- All code has been deposited and is publicly available on Github.					
18	Accession numbers are listed in the key resources table.					
19						
20	EXPERIMENTAL MODEL AND SUBJECT DETAILS					
21	Animals					
22	C57BI/6 mice were obtained from the Sanger Research support facility in-house-breeding					
23	colony. They were housed in a temperature and humidity-controlled facility in individually					
24	ventilated cages under a non-reversed light-dark cycle (Sanger Research support facility) or a					
25	reversed-light-dark cycle (ETH EPIC). Standard chow (LabDiet(r) 5021-3 supplied by IPS) and					

26 water were provided *ad libitum after weaning* unless stated otherwise (e.g. oocyte donors).

Breeding colony was provided SAFE R03-10 breeding diet, supplied by SAFE diets. Experimental procedures were performed during the animals' inactive cycle at Sanger. Age and weight matched (margin of one week) males were used in each experimental group receiving Dex injections. Animals used for Dex injection followed by sperm sequencing were all sexually mature (14 and 7 days or 3 hours post treatment were 13, 11 and 9 weeks of age respectively) at the time of sperm collection.

C57BI/6 males used for sperm sequencing 14 days post Dex injection and q-PCR
experiments/validation 3 hours post Dex were obtained from the ETH's EPIC in house
breeding colony in Zürich and were 14-18 weeks old. These mice were fed chow #3734 by
Kliba/Granovit.

11 IVF oocyte donor females and embryo recipients were fed SAFE R03-10 breeding diet, 12 supplied by SAFE diets until 10 days post embryo transfer. Until this time embryo recipients 13 were housed in pairs after which they were split into single housing. IVF offspring was weaned 14 at PND21 and assigned to cages avoiding littermate cohousing. Offspring phenotyping was 15 carried out between 3.5 to 4 months and necropsy at 4.5 months of age in balanced (offspring 16 controls, offspring treatment) and age matched groups (all animals had an age spread of 3 17 days). Animals were housed in groups of 4-5 mice/cage in the Sager Institute barrier research 18 support facility (all animals apart from animals for q-RT-PCR experiments) and ETHZ's EPIC 19 facility (animals for g-RT-PCR).

All experiments were approved by the UK home office (project license P176396F2) and Cantonal commission for animal experimentation Zürich (project license ZH222/19).

22

23 METHOD DETAILS

24 Dex treatment and sample collection

Age matched males with an age spread of 1 week were randomly assigned to control and treatment groups. Males were injected with either 2mg/kg of Dex in 10 % DMSO, 0.9% saline or vehicle (10%DMSO in 0.9% saline). Males used for sperm collection did not undergo any metabolic testing. They were sacrificed 2 weeks, 7 days and 3 hours after Dex or vehicle

treatment. Cauda epididymis and vas deference were dissected and placed in M2 medium.
After allowing sperm to diffuse into M2 medium, cells were pelleted by short centrifugation and
washed with PBS. For sperm RNA sequencing and q-PCR, mature sperm cells were separated
from potential somatic contamination by somatic lysis, followed by 2 washes with PBS
(Brykczynska *et al.*, 2010). Sperm counts and fertilization rate appeared unaffected post Dex
injection (Supplementary Fig. 6A,B).

7

8 In vitro fertilization and embryo culture

9 12 randomly selected, C57BL/6 females were superovulated at 26-31 days of age with Card
10 Hyperova (Cosmo Bio, KYD-010-EX-X5), followed by 7.5 IU human chorionic gonadotrophin
11 (HCG) 48 hours later.

12 Cumulus-oocyte complexes (COCs) were released from the ampulla of the oviduct 16-17 13 hours after HCG administration, and preincubated in high calcium HTF with Glutathione 14 medium for 30-60 minutes (in CO2 incubator at 37 deg C, 5% CO2 in air) before insemination. 15 Frozen sperm used for insemination was pooled from 2 males that had been injected with Dex 16 or vehicle 14 days prior to cryopreservation. Thawed sperm was preincubated for 30 minutes 17 in TYH (with Methyl-b-cyclodextrin, Sigma C4555) medium at 37 deg C, 5% CO2 in air, before 18 being added to the COC complexes for fertilisation. 4 hours after insemination, the presumptive 19 zygotes were washed through several drops of KSOM (Millipore, MR-121-D) and incubated 20 overnight in KSOM.

21 For in vivo offspring, 14-20 x 2 cell embryos from overnight culture in 6 individual IVF dishes 22 /group were implanted into 0.5 dpc pseudo-pregnant F1 females (6 females/group). Each dish 23 contained oocytes from one female with the exception of 2 dishes (out of 6) in the Dex group 24 that contained oocytes of the same female, since one female failed to super-ovulate. For 25 molecular (single) embryo gene expression analysis at the two cell stage, 2-cell-embryos from 26 overnight culture were frozen, and after thawing briefly cultured in preincubated KSOM 27 until/during plating into 96 well plates. The females used to generate these embryos were 28 superovulated with PMSG. The IVF Protocol is based on EMMA Harwell's protocol (adapted

from Takeo & Nakagata 2011(Nakagata, 2011)), and the Sperm Freeze Protocol is based on
 Ostermeier G.C. et al (2008)(Ostermeier *et al.*, 2008). Resulting litter sizes did not differ
 between Vehicle and Dex injected offspring (Supplementary Fig. 6C)

4

5 Sperm and embryo RNA extraction

Total RNA was prepared from adult mouse sperm using Trizol (Thermo Scientific 15596026)
and Directzol (Zymo R2080). Total RNA was prepared from zygotes using the Trizol LS
protocol. Quantity and purity of RNA were determined by Agilent 2100 Bioanalyser (Agilent
Technologies) and Qubit fluorometer (Life Technologies). Absence of prominent ribosomal
peaks indicated absence of somatic cell contamination.

11

12 Serum Exosome RNA extraction

13 Trunk blood was collected from animals following cervical dislocation and stored at room 14 temperature for 30 minutes to allow coagulation. Serum was subsequently separated by 2 15 centrifugation steps first for 10 min followed by 15 minutes at 3000. g. 100 ul of serum were 16 used as input for exosomal isolation following the manufacturers instructions (exoRNeasy 17 Qiagen). 2 ul of. 25 ul RNA eluate were used as input for cDNA conversion with the 18 miRCURY® LNA® cDNA conversion kit.

19

20

21 Sperm RNA sequencing (RNAseq)

Sequencing was done using an Illumina Genome Analyzer HiSeq 2500 (Illumina) in Rapid run
 mode for long 100bp and small 50 bp RNA sequencing runs respectively.

Libraries for long RNA sequencing were prepared using the TruSeq Stranded Total RNA kit according to the manufacturer's instructions with indices diluted at 1:3. 200 ng of total sperm RNA was subjected to removal of rRNA using Ribozero gold kit. Approximately 100ng of sperm RNA and total RNA of several 2-cell zygotes was subjected to TruSeq or Nextflex (sperm 14 days post injection) small RNA library preparation following the manufacturer's recommendations with the following modifications: adaptors were diluted 1:4 and PCR cycles
 were augmented to 18 and 22 (Nextflex) PCR cycles respectively. When library preparation of
 samples was split across days groups were balanced to circumvent batch effects.

4

5 Single embryo seq.

6 2 cell embryos were generated using the same conditions as indicated for in vivo offspring yet 7 followed by embryo cryopreservation until processing for library preparation. They were 8 thawed and those that appeared intact (34 controls and 37 Dex) pipetted into wells of 2 96 well 9 culture plates containing lysis buffer and stored at -80°C before processing according to the 10 Smartseq 2 protocol and manufacturer's recommendations (Nextera). Libraries contained a 11 1:19 Million dilution of External RNA Controls Consortium (ERCC) spike-ins (4456740 Ambion) 12 and were amplified for 18 PCR cycles. Sequencing was performed on a HiSeq V4 under paired 13 end 75bp mode.

14

15 Insulin and Glucose tolerance test

Animals were fasted 4 hours to establish a shared baseline glucose level. They received a single injection of insulin (insulin: 1mU/g body weight) (Actrapid Novo Nordisk), glucose (2mg/g body weight) or vehicle (saline) intraperitoneally. Blood samples were taken from lateral tail vein in adult animals to assess blood glucose level using an Accuckeck aviva device.

20

21 Body mass index

22 Animal lengths were measured using a standard ruler and weighed for assessing body weight.

23 Body mass index was calculated using the following formula: weight (g)/(length (cm)^2).

24

25 Necropsy

Organs were dissected after sacrifice and weighed immediately on a scale using "g" as a unit
with an accuracy of 2 decimals (accurate down to 10 mg).

1 Small RNA q-RT-PCR

5ng/sample RNA isolated from sperm was reverse transcribed (RT) using the miCURY LNA RT kit (Qiagen #339340). Quantitative RT-PCR (qRT-PCR) was performed using SYBR green based detection in a Biorad thermal cycler with MiRCURY LNA-based small RNA probes designed against tRNA ArgCCT-2, Gly-GCC-2, His-GTG-2, Leu-CAA-2, with a polyA tail directed reverse miRCURY primer (Qiagen # 339317). RnU6 was used as an internal control in sperm samples and mir-103a-3p in serum samples (Qiagen # 339306).

8

9 CircRNA q-RT-PCR

100ng/sample RNA isolated from sperm was converted into cDNA using random hexamers.
Primers were designed to span the exon splice junctions. Primer sequences for CircTasp1 and
CircDennd1b are depicted in the resource table. Tubulin1 was used as endogenous control in
the Sybr Green based quantification.

14

15 QUANTIFICATION AND STATISTICAL ANALYSIS

16 Bioinformatic analysis

17 Bulk RNA sequencing

18 Each sequencing library represented sperm harvested from a single male. Sequencing 19 quality was assessed with FastQC (Andrews, 2010) and MultiQC (Ewels et al., 2016). 20 Adapters were removed from the 3' ends with cutadapt (Martin, 2011) (version 1.14) and 21 resulting sequences with 14 nucleotides of length or less were discarded. All other reads 22 were aligned end to end (no soft clipping) to the ENSEMBL *Mus musculus* genome (release 23 75) (Flicek et al., 2014) with STAR (Dobin et al., 2013). No mismatches were allowed. 24 Featurecounts was used to match the alignments against the miRbase(Kozomara and 25 Griffiths-Jones, 2011) annotation (version 21) and obtain a matrix of miRNA counts. We 26 applied fractional counts whenever alignment occurred at multiple genomic locations. 27 Differential expression was analyzed using DESeq2 (Love, Huber and Anders, 2014). 28 Quantification of tRNA fragments was performed as above, but all CCA-3' trinucleotides were trimmed after adapter removal, sequences with 15 nucleotides or less were subsequently
 discarded and GtRNAdb (Chan and Lowe, 2016) annotation (GRCm38/mm10) was used to
 obtain the count matrix. Quantification of rRNA fragments was performed using SPORTS(Shi
 et al., 2018) on the precompiled database included in the tool.

5

For the data set collected 14 days after Dex injection, library preparation included the insertion
of 2 random tetranucleotides between read and adapters. By including only unique sequences
in the analysis we removed duplicates due to PCR amplification.

9 Long RNAseq libraries were pre-processed with trimmomatic (Bolger, Lohse and Usadel,

2014) to remove adapters. Reads were aligned to the genome using STAR (Dobin *et al.*, 2013) and quantified using featurecounts (Liao, Smyth and Shi, 2014). Circular RNAs were quantified using Circexplorer2 (Zhang *et al.*, 2016) based on junction reads as detected by STAR. Differential expression analysis was performed on the combined set of counts for circular and non-circular RNAs using edgeR (Anders *et al.*, 2013). Robust estimation of dispersion was used to avoid spurious significance due to outliers.

16

17

2-cell single embryo sequencing analysis

18 Reads from 2 cell embryos were mapped to the mouse reference genome (mm10) and ERCC 19 spike-ins using STAR (Dobin et al., 2013). Resultant alignments were processed to quantify 20 the expression of annotated genes by GENCODE (vM11) and ERCC spike-ins using 21 featureCounts (Liao, Smyth and Shi, 2014). To filter low-guality sequenced embryos we only 22 considered those which had a total read count of at least 0.5 million reads with less than 15% 23 and 10% their read counts mapping to mitochondrial genes and ERCC spike-ins respectively. 24 After these filters were applied a total of 56 embryos (29 controls and 27 treated) remained. 25 We clustered their gene expression profiles using SC3 (Kiselev et al., 2017) obtaining two main

1 clusters (C1 and C2). Using scmap (Kiselev, Yiu and Hemberg, 2018), we projected the gene 2 expression profiles for the two cell embryos onto an index containing expression profiles from 3 zygotic, early/mid/late 2 cell embryos and 4 cell embryo cells reported by Deng et al. (Deng et 4 al., 2014). We performed PCA analyses using scater (McCarthy et al., 2017) (runPCA function) 5 and we calculated the PC1 silhouette coefficient using in-house R scripts. To perform 6 differential gene expression analyses we normalized the read counts of each embryo as FPKM 7 and we used Census (Qiu et al., 2017) algorithm to convert these values into relative transcripts 8 counts. We computed the obtained 'Census counts' using Monocle (v 2.99.2), assuming a 9 negative binomial distribution and a lower detection limit of 0.5. We performed differential gene 10 expression analyses between the total treated and control embryos, and also between the 11 treated and control embryos inside of C1 and C2 clusters.

12

13 Remaining statistical analyses.

14 Sample size for in vivo offspring phenotyping was estimated based on previous work on similar 15 models (Hoyer et al., 2013; Bohacek, von Werdt and Mansuy, 2016). 3-Way repeated measures ANOVA was used to assess statistical significance for BMI, GTT and ITT 16 17 measurements. Necropsy data were analysed using 3-way ANOVA followed by multiple t-tests 18 and corrected for multiple comparisons using the Benjamini-Hochberg method. Normality was 19 assessed with the Kolmogorov Smirnov test and met in all necropsy data. Homogeneity of 20 variances was assessed and met in all necropsy data unless gonadal WAT. These t-tests did 21 not assume homogeneity of variances (applied Welchs correction). Q-RT-PCR results 22 comparing caput and cauda sperm RNA were analyzed by fitting a Mixed model followed by 23 posthoc tests to compare individual groups applying the Sidak correction for multiple 24 comparisons. Other q-PCRs were analysed using multiple t-tests corrected for multiple 25 comparisons applying the two stage step up method by Benjamini, Krieger and Yekutieli. All 26 reported replicates were biological replicates. Significance was set at p < 0.05 and where 27 applicable q<0.05 for all tests. All statistics of behavioural, metabolic tests and q-RT-PCR were

1	computed with Prism. Outliers were removed from q-PCR results using Prism^s inbuild ROUT
2	method and are depicted in the supplementary tables containing raw data with a star.
3	
4	
5	Supplementary Material
6	This article contains supplementary Figures and tables. Supplementary figures and legends
7	are compiled in one document.
8	
9	Supplementary table legend
10	
11	Supplementary Table 1 related to Figure 1: Data and analysis of sperm small RNA 14 days
12	post Dex
13	Sheet 1: List of normalized miRNA read counts of sperm harvested 14 days post Dex and
14	vehicle injection.
15	Sheet 2: Deseq2 results of a comparison between miRNA from sperm harvested 14 days
16	post Dex and vehicle injection.
17	Sheet 3: List of normalized tsRNA read counts of sperm harvested 14 days post Dex and
18	vehicle injection.
19	Sheet 4: Deseq2 results of a comparison between tsRNAs from sperm harvested 14 days
20	post Dex and vehicle injection.
21	Sheet 5: List of normalized read counts for rRNA from sperm harvested 14 days post Dex
22	and vehicle injection.
23	Sheet 6: Deseq2 results for rRNA quantification from sperm harvested 14 days post Dex and
24	vehicle injection.
25	Sheet 8: Raw values of qRT-PCR analysis for serum.
26	Sheet 9: Raw values of qRT-PCR analysis of caput sperm.
27	
28	

1 Supplementary table 2 related to Figure 2: Data and analysis of sperm small RNA 3 hours

2 and 7 days post Dex

3 Sheet 1: Deseq2 results of the comparison of tsRNAs in sperm harvested 3 hours and 7

4 days post Dex with tsRNAs in sperm harvested 3 hours and 7 days post vehicle injection.

5 Sheet 2: Deseq2 results of the comparison between tsRNAs from sperm harvested post Dex

6 and vehicle injection at 3 hours with tsRNAs in sperm harvested 7 days post Dex and vehicle

7 injection.

8 Sheet 3: Deseq2 results of the interaction between treatment and time for tsRNAs from

9 sperm harvested 3 hours and 7 days post Dex and vehicle injection.

10 Sheet 4: Deseq2 results of the comparison of miRNAs in sperm harvested 3 hours and 7

11 days post Dex with miRNAs in sperm harvested 3 hours and 7 days post vehicle injection.

12 Sheet 5: Deseq2 results of the comparison between miRNAs from sperm harvested post Dex

13 and vehicle injection at 3 hours with miRNAs in sperm harvested 7 days post Dex and

14 vehicle injection.

15 Sheet 6: Deseq2 results of the interaction between treatment and time for miRNAs from

16 sperm harvested 3 hours and 7 days post Dex and vehicle injection.

17 Sheet 7: Raw values of qRT-PCR analysis for tsRNA-Arg-CCT-2 in caput and cauda sperm

18 sampled 3 hours post Dex and vehicle injection.

19

20 Supplementary table 3 related to Figure 3: Raw data for Dex offspring phenotyping.

21 Raw data for BMI (sheet 1), GTT (sheet 2) , ITT (sheet 3) and necropsy weights (sheet 4) of

22 adult offspring animals resulting from IVF of wildtype oocytes and sperm harvested 14 days

23 post Dex and vehicle injection.

24

25 Supplementary table 4 related to Figure 5: Single embryo RNA sequencing analysis.

26 Sheet 1: Monocle output list of significantly differentially regulated genes between 2-cell

27 embryos resulting from IVF of wildtype oocytes and sperm harvested 14 days post Dex and

28 vehicle injection.

38

3	post Dex	
4		
5	Sheet 1: List of normalized long RNA seq counts of sperm harvested 14 days post Dex and	
6	vehicle injection.	
7	Sheet 2: EdgeR results of a comparison between long RNA reads from sperm harvested 14	ł
8	days post Dex and vehicle injection.	
9	Sheet 3: Raw values of qRT-PCR analysis for caput and cauda sperm circRNAs.	
10		
11		
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Supplementary table 5 related to Figure 6: Data and analysis of sperm long RNA 14 days

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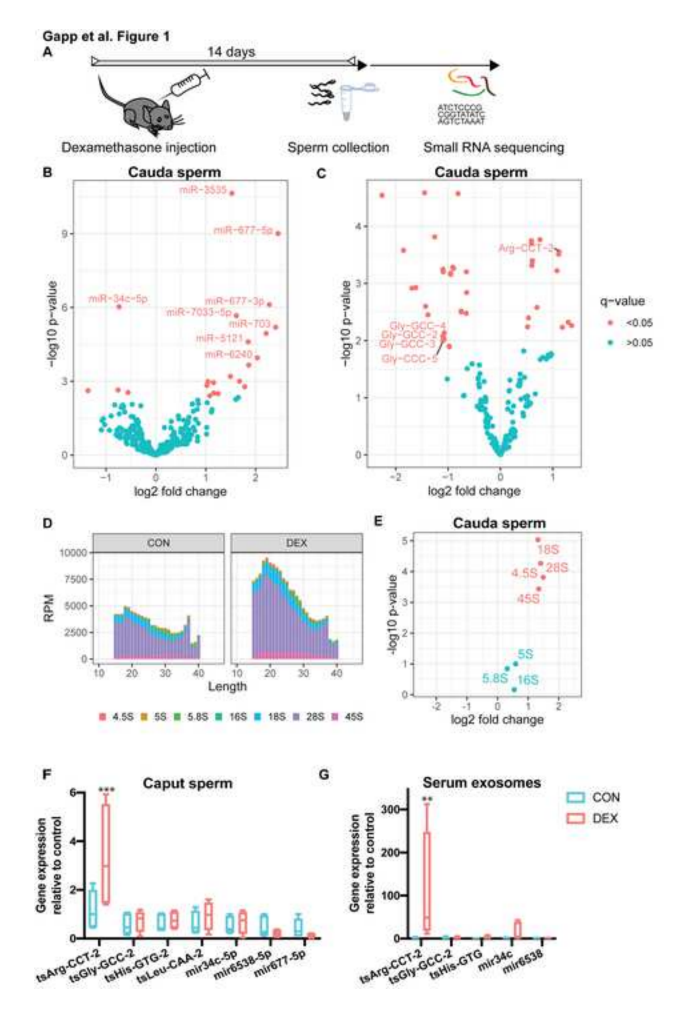
CellPress

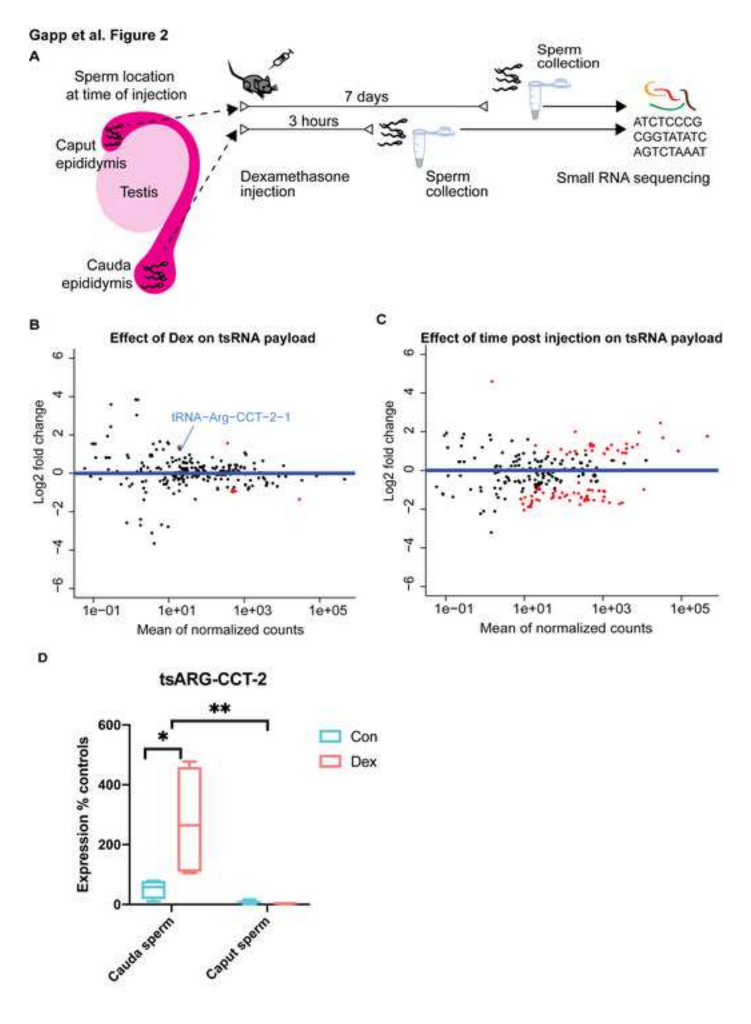
KEY RESOURCES TABLE

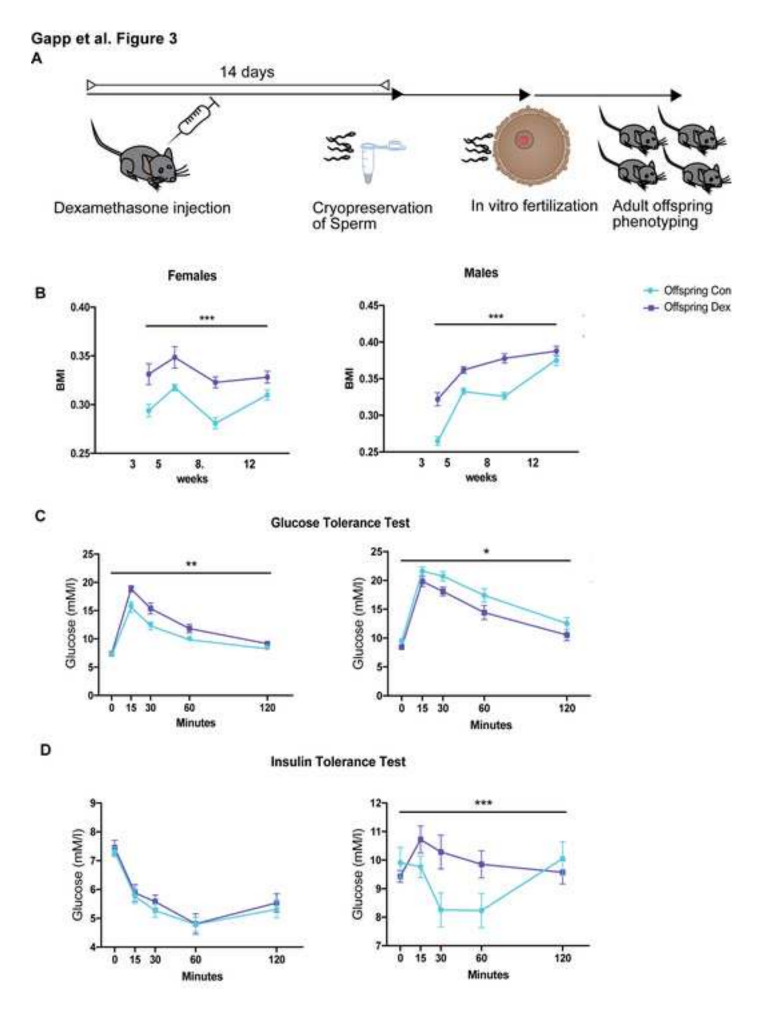
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, peptides, and recombinant proteins		
Dexamethasone	Sigma	Cat#D4902-25MG
DMSO	VWR	Cat#472301- 100ML
Hyperova	Cosmo Bio	Cat#KYD-010-EX- X5
KSOM	Millipore	Cat#MR-121-D
Trizol	Thermo Scientific	Cat#15596026
Directzol	Zymo	Cat#R2080
M-MLV	Promega	Cat#M1701
Recombinant RNAsin Ribonuclease inhibitor	Promega	Cat#N2111
Insulin	Novo Nordisk	Cat#Actrapid
Critical commercial assays		
exoRNeasy Serum/Plasma Midi Kit	Qiagen	Cat#77044
Truseq small RNA library kit	Ilumina	Cat#RS-200-0012
Truseq Total RNA library kit	Ilumina	Cat#RS-122-2301
Nextflex small RNA library kit	Perkin Elmer	Cat#NOVA-5132- 05
Nextera XT DNA Library Preparation Kit	Ilumina	Cat#FC-131-1024
miRCURY LNA RT Kit	Qiagen	Cat#339340
miRCURY® LNA® miRNA SYBR® Green PCR	Qiagen	Cat#339345
Deposited data		
Sperm small and long RNA sequencing data	Gene Omnibus	GSE162112
2-cell embryo RNA sequencing data	ENA	ERP105660
Experimental models: Organisms/strains		
C57BI/6Jrj mice	Janvier lab	
C57BI/6 CBLT mice	Sanger Institute	
Oligonucleotides		
mirCURY LNA miRNA PCR Assay ArgCCT-2 5`GCCCCAGUGGCCUAAUGGAUAAGGCACUGGCC3`	Qiagen	Cat#339317
mirCURY LNA miRNA PCR Assay Gly-GCC-2 5`GCAUUGGUGGUUCAGUGGUAGAAUUCUCGCCU3 `	Qiagen	Cat#339317
mirCURY LNA miRNA PCR Assay His-GTG-3 5`GCCGUGAUCGUAUAGUGGUUAGUACUCUGCGU3	Qiagen	Cat#339317
mirCURY LNA miRNA PCR Assay Leu-CAA-2 5`GUCAGGAUGGCCGAGUGGUCUAAGGCGCCAGA3	Qiagen	Cat#339317
mirCURY LNA miRNA PCR Assay RnU6	Qiagen	Cat#339317
mmu-miR-677-p5 mirCURY LNA miRNA PCR Assay	Qiagen	Cat#339306
mmu-miR-3535 mirCURY LNA miRNA PCR Assay	Qiagen	Cat#339306
mmu-miR-6538 mirCURY LNA miRNA PCR Assay	Qiagen	Cat#339306
CircTasp1 FW `CTT AGG AGA GAT TGA ATG TGA TGC C` RW `AAA GGG AGT CAA CCA CTC AG`	Microsynth	Cat#4059186 & 4059187



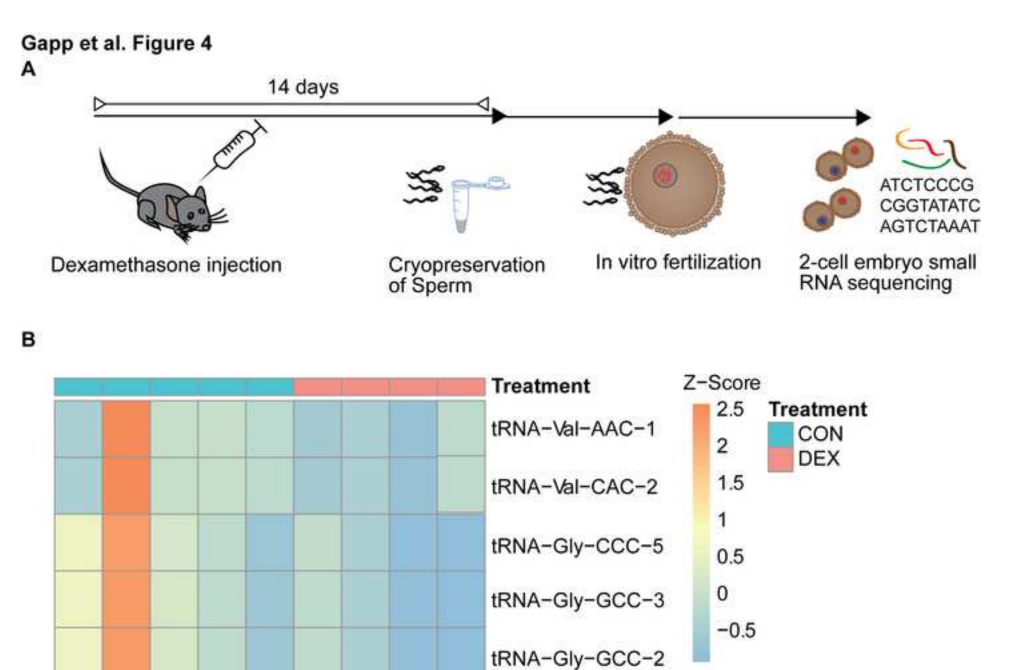
CircDennd1 FW `AGCTTTCCCAGTTTATTGATGGT` RW `GAAGCCACCCGAAGTGATCT`	Microsynth	Cat#4059182 & 4059183
Software and algorithms		
Code	Github	https://github.com/ ETHZ-INS/Sperm- RNA-Dex).
GraphPad Prism	GraphPad Prism version 8	www.graphpad.co m
featureCounts	Liao et al., 2014	http://subread.sou rceforge.net/
DESeq2	Love et al., 2014	https://bioconduct or.org/packages/r elease/bioc/html/D ESeq2.html
edgeR	Anders <i>et al.</i> , 2013	https://bioconduct or.org/packages/r elease/bioc/html/e dgeR.html
STAR	Dobin et al., 2013	https://github.com/ alexdobin/STAR
CIRCexplorer2	Zhang et al., 2016	https://circexplorer 2.readthedocs.io
scater	McCarthy et al., 2017	https://bioconduct or.org/packages/r elease/bioc/html/s cater.html
Monocle	Qiu et al., 2017	http://cole- trapnell- lab.github.io/mono cle-release/
Trimmomatic	Bolger et al., 2014	www.usadellab.or
cutadapt	Martin, 2011	https://cutadapt.re adthedocs.io



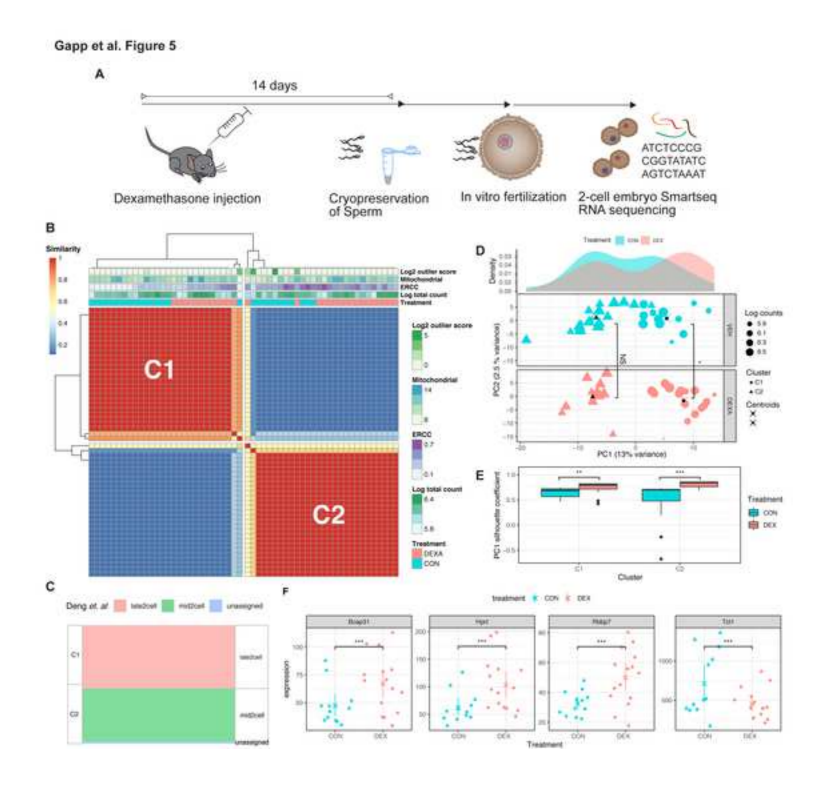


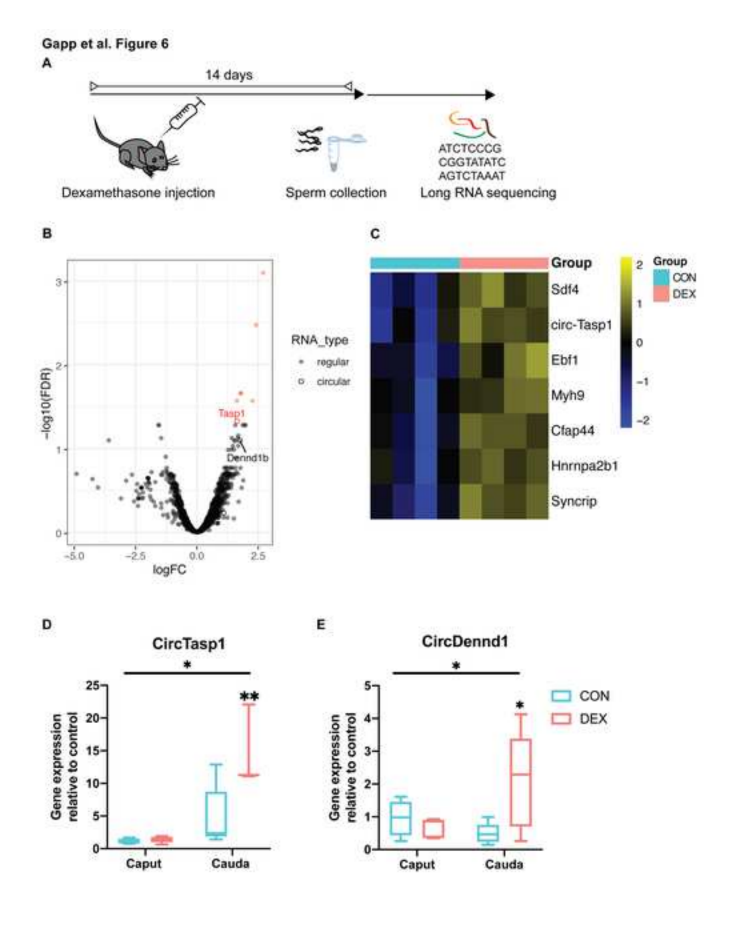


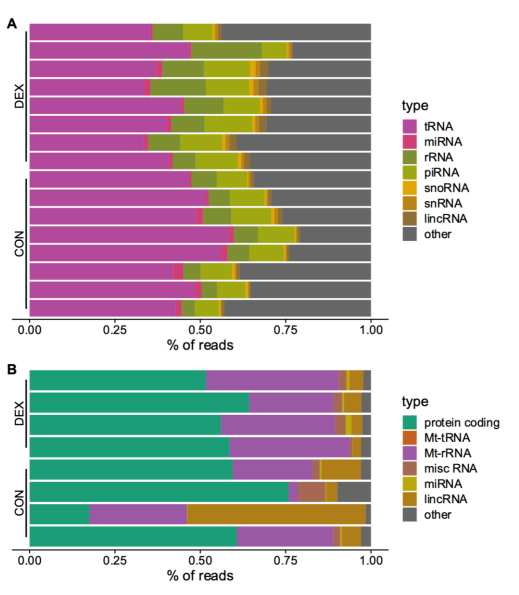




tRNA-Gly-GCC-4





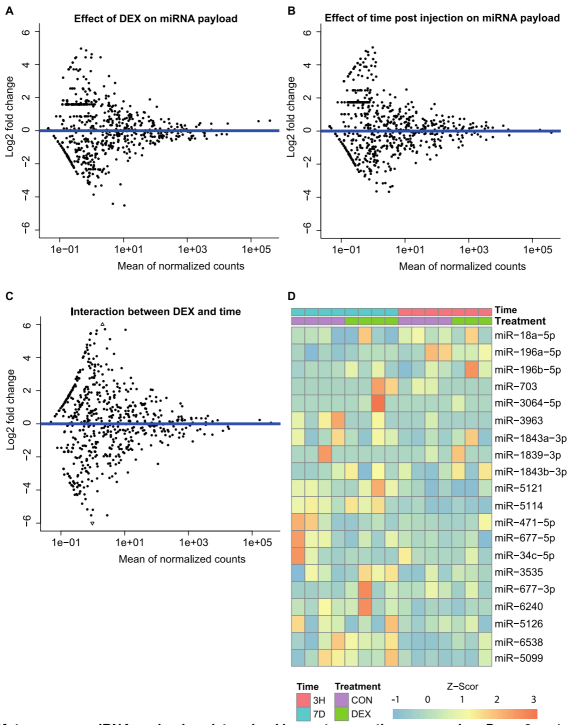


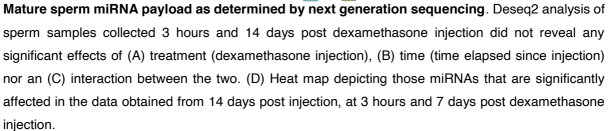
Supplementary Information

Supplementary Figure 1 related to Figure 1 and 4

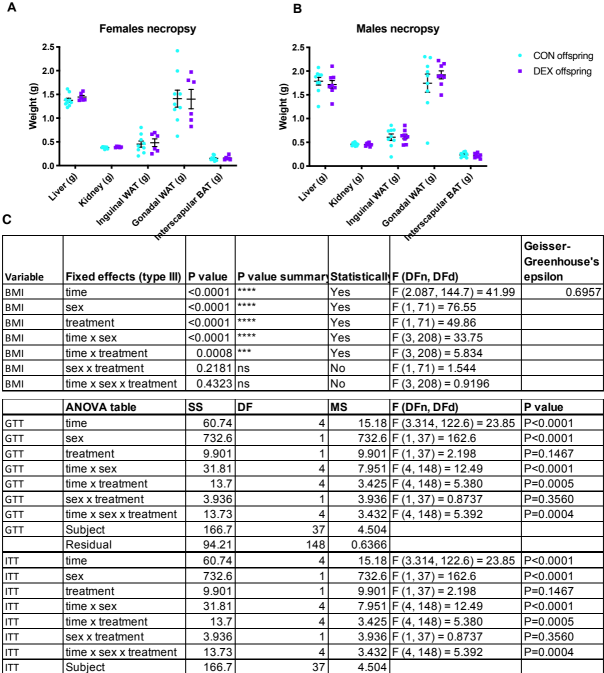
Relative percentage of RNA Ensemble biotypes and tRNA matching reads (A) in small RNA sequencing libraries of 8 sperm samples from control males and 8 sperm samples 14 days post dexamethasone injection. (B) long RNA sequencing libraries of 4 sperm samples from control males and 4 sperm samples 14 days post dexamethasone injection.

Supplementary Figure 2 related to Figure 2





Supplementary Figure 3 related to Figure 3



0.6366

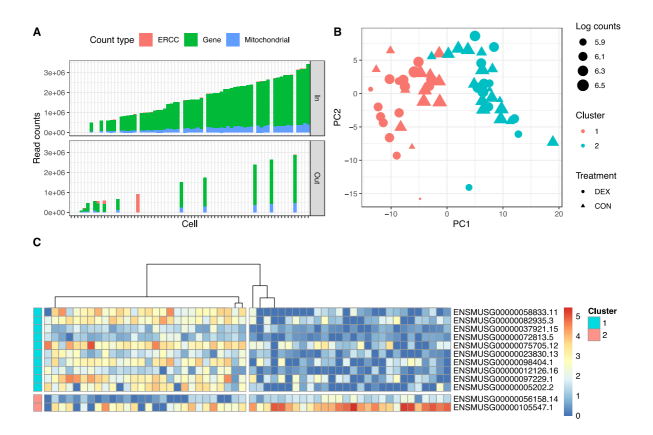
148

Tissue	P value	Difference	SE of difference	t ratio
Liver (g)	0.27665	-0.06789	0.05979	1.136
Kidney (g)	0.188702	-0.01267	0.009131	1.387
Inguinal WAT (g)	0.776367	-0.02972	0.1025	0.29
Gonadal WAT (g)	0.968101	0.01122	0.2753	0.04077
Interscapular BAT (g)	0.801259	-0.006444	0.02508	0.2569
Liver (g)	0.562456	0.06872	0.116	0.5923
Kidney (g)	0.788758	0.004292	0.01573	0.2728
Inguinal WAT (g)	0.872972	-0.014	0.08608	0.1626
Gonadal WAT (g)	0.434173	-0.1747	0.2174	0.8036
Interscapular BAT (g)	0.432084	0.01892	0.02343	0.8073

94.21

Residual

Necropsy followed by weighing of dissected tissue from offspring of control injected and Dexamethasone injected males. Comparison of the weight of liver, kidney, inguinal and gonadal white adipose tissue (WAT) as well as interscapular brown adipose tissue (BAT) did not reveal significant differences between (A) female and (B) male offspring of control and dexamethasone injected males. (C) Statistical values obtained by overall ANOVAs (BMI, GTT, ITT) and Multiple t-tests corrected for multiple comparisons (necropsy). All Data besides male gonadal WAT showed equal distribution of variances. Graphs show scattered dot plots with standard error of the mean. ITT = insulin tolerance test, GTT= Glucose tolerance test, BMI= Body mass index, WAT = white adipose tissue. BAT= brown adipose tissue.



Supplementary Figure 4 related to Figure 5

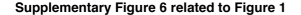
Single- 2-cell embryo sequencing using the Smartseq method. (A) Result of the quality filters implemented to select the 2 cell embryos that were used for downstream analyses. Libraries (embryos/cells) that contained a suboptimal number of mitochondrial (>15%) or ERCC mapping reads (>10%) or yielded less than 500 000 reads were excluded (displayed in Out). Most libraries showed a high number of gene mapping reads and were retained and processed for further analysis (displayed in In). (B) PCA results shown as a single panel, revealing segregation of embryos for the first two principle components (matching almost perfectly cluster 1 (red) and cluster 2(blue)) and additionally depicting read count number (size of circle) and assignment to treatments (offspring embryos of control fathers = CON, dots and fathers who were injected with dexamethasone 14 days prior sperm harvest = DEX, triangles). PC2 is attributed to technical factors such as read counts. (C) Marker genes identified by sc3 for C1 and C2 clusters.

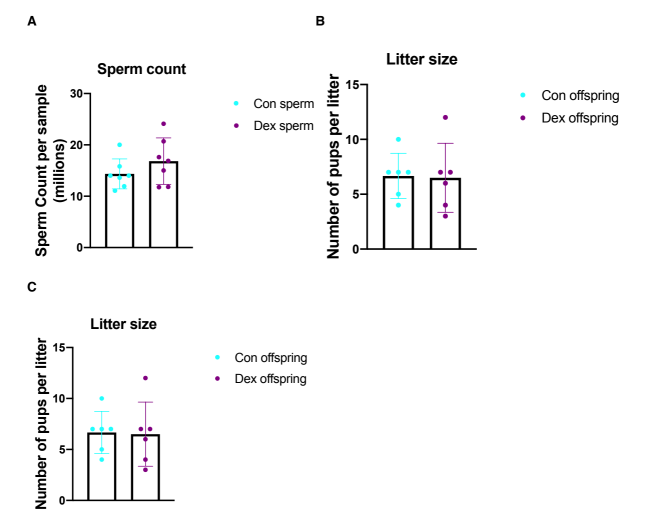
Supplementary Figure 5 related to Figure 5 and 6

circAltas ID	microRNA name	#binding sites of miRanda	#bingding sites of targetScan
mmu-Dennd1b_0017	<u>mmu-miR-201-3p</u>	1	1
mmu-Dennd1b_0017	<u>mmu-miR-3110-5p</u>	1	1
mmu-Dennd1b_0017	<u>mmu-miR-1953</u>	1	1
mmu-Dennd1b_0017	<u>mmu-miR-1981-5p</u>	1	1
mmu-Dennd1b_0017	<u>mmu-miR-706</u>	1	1
mmu-Dennd1b_0017	<u>mmu-miR-6363</u>	1	1
mmu-Dennd1b_0017	mmu-miR-3074-5p	1	1
mmu-Dennd1b_0017	mmu-miR-1955-5p	1	2
mmu-Dennd1b 0017	mmu-miR-675-3p	1	1
mmu-Dennd1b_0017	mmu-miR-343	1	2
mmu-Dennd1b_0017	mmu-miR-3083-5p	1	1
mmu-Dennd1b_0017	mmu-miR-3073a-5p	1	1
mmu-Dennd1b_0017	mmu-miR-9-5p	1	1
mmu-Dennd1b_0017	mmu-miR-378b	1	1
<u>mmu-Tasp1_0039</u>	<u>mmu-miR-221-5p</u>	1	1
mmu-Tasp1_0039	mmu-miR-5627-3p	1	3
mmu-Tasp1_0039	<u>mmu-miR-298-5p</u>	1	3
mmu-Tasp1_0039	mmu-miR-3058-5p	1	1
mmu-Tasp1_0039	<u>mmu-miR-207</u>	1	3
mmu-Tasp1_0039	<u>mmu-miR-709</u>	1	1
mmu-Tasp1_0039	mmu-miR-6537-3p	1	1
mmu-Tasp1 0039	mmu-miR-713	1	2
mmu-Tasp1_0039	mmu-miR-3094-3p	1	2
mmu-Tasp1_0039	mmu-miR-100-3p	1	2
mmu-Tasp1 0039	mmu-miR-1906	1	3
mmu-Tasp1 0039	mmu-miR-320-5p	1	3
mmu-Tasp1 0039	mmu-miR-3082-3p	1	2
mmu-Tasp1_0039	mmu-miR-706	1	1
mmu-Tasp1_0039	mmu-miR-3110-5p	1	1
mmu-Tasp1_0039	mmu-miR-6900-5p	1	1
mmu-Tasp1_0039	<u>mmu-miR-1955-5p</u>	1	-
mmu-Tasp1_0039	mmu-miR-674-5p	1	1
mmu-Tasp1_0039	mmu-miR-351-3p	1	1
mmu-Tasp1_0039	<u>mmu-miR-5104</u>	1	1
mmu-Tasp1_0039	mmu-miR-504-3p	1	1
mmu-Tasp1_0039	mmu-miR-152-5p	1	1
mmu-Tasp1_0039	mmu-miR-1897-5p	1	1
mmu-Tasp1_0039	mmu-miR-3105-5p	1	1
mmu-Tasp1_0039	<u>mmu-miR-742-3p</u>	1	2
mmu-Tasp1_0039	mmu-miR-1969	1	2
mmu-Tasp1_0039	mmu-miR-1960	1	2
<u>mmu-Tasp1_0039</u>	mmu-miR-24-3p	1	2
mmu-Tasp1_0039	mmu-miR-188-5p	1	1

CircRNA targets as predicted by CircAtlas. Each column displays results from miRanda and targetScan.

miRNAs that are targeted by both circRNAs are highlighted in purple.





Fertility measures following Dexamethasone treatment. (A) Sperm cell count 14 days post vehicle (n=7) or Dex (n=7) injection did not reveal any difference in sperm number (t(12)=1.221, p=0.25) suggesting no impairing effect of Dex on spermatogenesis. (B) Count of fertilized oocytes (as of the appearance of the second pronucleus) over total available oocytes of 6 replicates of cryopreserved sperm from a pool of 2 dex injected versus 2 vehicle injected males did reveal no significant difference in the fertilization rate between dex and vehicle sperm. (C) Number of pups per litter was similar in offspring resulting from in vitro fertilization with sperm from fathers that were injected with Vehicle (n=6) or Dex (n=6) (t(12)=1.122, p=0.25).