Genetically-biased fertilization in APOBEC1 complementation factor (A1cf) mutant mice

Naoki Hirose1, Genevieve Blanchet1, Yasuhiro Yamauchi1, Abigail C. Snow2, Robin Friedman3,5, Carmen Y. Khoo4, Christine W. Lary4, Monika A. Ward1,6,7 & Joseph H. Nadeau2,6,7

Meiosis, recombination, and gametogenesis normally ensure that gametes combine randomly. But in exceptional cases, fertilization depends on the genetics of gametes from both females and males. A key question is whether their non-random union results from factors intrinsic to oocytes and sperm, or from their interactions with conditions in the reproductive tracts. To address this question, we used in vitro fertilization (IVF) with a mutant and wild-type allele of the A1cf (APOBEC1 complementation factor) gene in mice that are otherwise genetically identical. We observed strong distortion in favor of mutant heterozygotes showing that bias depends on the genetics of oocyte and sperm, and that any environmental input is modest. To search for the potential mechanism of the ‘biased fertilization’, we analyzed the existing transcriptome data and demonstrated that localization of A1cf transcripts and its candidate mRNA targets is restricted to the spermatids in which they originate, and that these transcripts are enriched for functions related to meiosis, fertilization, RNA stability, translation, and mitochondria. We propose that failure to sequester mRNA targets in A1cf mutant heterozygotes leads to functional differences among spermatids, thereby providing an opportunity for selection among haploid gametes. The study adds to the understanding of the gamete interaction at fertilization.

Discovery that bias is evident with IVF provides a new venue for future explorations of preference among genetically distinct gametes at fertilization for A1cf and other genes that display significant departure of Mendelian inheritance.

The random nature of fertilization seemed so obvious that Mendel did not include it in his Laws, even though it is essential for understanding the inheritance of phenotypic variation. Mendel’s appreciation for the fundamental role of chance in sexual reproduction is evident in his use of binomial probabilities to convert frequencies of haploid gametes (A or a) into frequencies of diploid individuals (AA, 2Aa, aa)1. To ensure Mendelian inheritance, haploid gametes in each sex must be functionally equivalent2. The strong selective pressure to maintain equal transmission limits phenotypic variation during meiosis, among gametes, and at fertilization3,4. Cytoplasmic bridges promote phenotypic similarity by enabling transfer of RNAs, proteins and other cellular constituents among the four genetically unique products of each meiosis5–7. However, some gene products are found primarily in the gamete in which they originate. This restricted localization can lead to phenotypic variation among haploid gametes8–11, creating opportunities for preferential transmission, with several of these genes carrying signatures of positive natural selection12.

Some genetic variants violate the Law of Independent Segregation by preferentially transmitting alleles from heterozygotes to offspring3,13–15. Several kinds of distorted transmission have been reported. Meiotic drive results from preferential segregation of alleles through meiosis. Transmission ratio distortion results from differences in gamete numbers and activity. Deviations from Mendelian expectations can also result from selective embryo, neonatal, or juvenile loss. Distortion is usually a property of one sex and is independent of the genetic constitution of their mating partner3,4,13. Although generally rare, examples of transmission distortion have been reported in many species3,13–15.

1Institute for Biogenesis Research, John A. Burns School of Medicine, University of Hawaii at Manoa, Honolulu, HI, USA. 2Center for Molecular Medicine, Maine Medical Center Research Institute, Portland, ME, USA. 3Ohana Biosciences, Cambridge, MA, USA. 4Center for Outcomes Research, Maine Medical Center Research Institute, Portland, ME, USA. 5Present address: Dragonfly Therapeutics, Waltham, MA, USA. 6These authors contributed equally: Monika A. Ward and Joseph H. Nadeau. &email: mward@hawaii.edu; joseph.nadeau@mainehealth.org
Discovery that fertilization can depend on the genetic constitution of both oocyte and sperm highlights a largely unexplored dimension of haploid selection. For example, among offspring of mouse crosses involving an engineered mutation in the *A1cf* gene (APOBEC1 complementation factor), a strong excess of heterozygotes was observed, as high as ~13 mutant heterozygotes found for every wild-type homozygote, instead of the 2:1 ratio expected for Mendelian inheritance in intercrosses (Fig. 1A). At least eight other genes similarly affect fertilization, with some yielding an excess of heterozygotes and others a deficiency. For these nine genes, mutant homozygotes are invariably absent.

**Figure 1.** Mendelian segregation and deviations from expectations. (A) Mendelian segregation. ‘m’ and ‘+’ designate the mutant and wild-type allele, respectively. Intercrosses between inbred strains make F1 hybrids. Hybrids are then crossed to either parental strain for backcrosses or *inter se* for intercrosses. Genotype ratios are based on Mendelian expectations. (B) Distinct segregation ratios, litter size, and genotype loss depending on the basis for non-Mendelian segregation. With embryonic lethality, mutant alleles that compromise viability of mm homozygotes can also result in loss of some m+ heterozygotes (partial dominance), which can further reduce litter size. Intercrosses can show an excess (<1 : >2) or a deficiency (1:1) of m+ relative to ++.

| Source of distortion | Cross      | Segregation (++ : m+ : mm) | Litter size | Genotype loss |
|----------------------|------------|----------------------------|-------------|---------------|
| Embryonic Lethality  | Backcross  | 1 : 1                      | Normal      | None          |
|                      | Intercross | 1 : 2 : 0                  | Reduced     | None          |
|                      | Intercross | 1 : <2 : 0                 | Reduced     | mm, some m+   |
| Distorted Transmission| Backcross  | <1 : >1                    | Normal      | None          |
|                      | Intercross | <1 : >2 : 0                | Reduced     | mm            |
| Biased Fertilization | Backcross  | 1 : 1                      | Normal      | None          |
|                      | Intercross | <1 : >2 : 0                | Normal      | None          |
|                      | Intercross | 1 : 1 : 0                  | Normal      | None          |
and Methods), the possibility of microbiome differences in the reproductive tracts between sites32–34, undetected source (mice derived recently from frozen embryos at UH versus a longstanding colony at MMCRI; see Materials
The somewhat variable magnitude of bias between UH and MMCRI may be explained by the differences in mouse
expectations or on the observed backcross results (Text S1). Backcrosses at MMCRI but not UH showed modest
offspring (6:1) in the MMCRI intercross and 3.1 m+ for every ++ offspring (3:1) in the UH intercross, instead of
backcrossed to ++ homozygotes (Fig. 2). These breeding experiments were performed at two locations, the Maine
Medical Center Research Institute (MMCRI) and the University of Hawaii (UH). We found 6.1 m+ for every ++
reproductive features to test alternative explanations for bias and mined available transcriptomes to identify
factors intrinsic to gametes from epididymal sperm, from A1cf
A1cf
influence of somatic effects from the parents on segregation, we obtained gametes, MII oocytes and cauda epididymal sperm, from A1cf
in the oviduct25,26, genetic interactions between factors in eggs and sperm during completion of meiosis at fertilization23,24, or from somatic factors that facilitate fertilization in the oviduct29–31. Here, we used in vitro fertilization (IVF) to test for gamete- versus soma-specific control of biased fertilization. We also measured reproductive features to test alternative explanations for bias and mined available transcriptomes to identify targets of the A1CF RNA-binding protein that could drive haploid selection. We demonstrated that the fertilization bias specifically depends on the genetics of oocyte and sperm and propose that it originates from the functional differences between spermatids arising from sequestration of A1cf transcripts. Previously, evidence for bias was incidental to the primary focus of reported work. Here, we report the first study focused exclusively on the evidence for bias and tests for one of the proposed mechanisms.

Results
In vitro fertilization and in vivo validation support unusual A1cf segregation. To mitigate the influence of somatic effects from the parents on A1cf segregation, we obtained gametes, MII oocytes and cauda epididymal sperm, from A1cf
parents and subjected them to in vitro fertilization (IVF). We examined a total of 332 IVF-produced embryos and a total of 1,198 mice resulting from various intercrosses (test) and backcrosses (control). To assess the relative contribution of gametes and soma to fertilization bias in vitro crosses were made with an appropriate combination of oocytes and sperm from A1cf
(mutant heterozygotes and A1cf
(++) wild-type homzygotes (Table 1, top). The ratio of m+ to ++ IVF-derived embryos (10:1) differed significantly from the expected 2:1 Mendelian ratio (χ² = 110.46, p < 0.0001). By contrast, backcross IVF results were consistent with expectations. Embryos derived from these crosses progressed similarly through early developmental stages in vitro (Table 1, bottom). The significant departure from expectations shows that bias depends largely on factors intrinsic to gametes from A1cf
heterozygotes, or perhaps on a genetically determined predisposition of gametes to somatic influences taking place during epididymal sperm maturation and oviductal oocyte transport persisting after gametes are isolated for IVF tests.

To verify bias in vivo, A1cf
heterozygotes were intercrossed, and in parallel m+ females and males were backcrossed to ++ homozygotes (Fig. 2). These breeding experiments were performed at two locations, the Maine Medical Center Research Institute ( MMCRI) and the University of Hawaii (UH). We found 6.1 m+ for every ++ offspring (6:1) in the MMCRI intercross and 3.1 m+ for every ++ offspring (3:1) in the UH intercross, instead of the 2:1 ratio expected for intercrosses. The intercross results were significant when based either on Mendelian expectations or on the observed backcross results (Text S1). Backcrosses at MMCRI but not UH showed modest departures from expectations (Fig. 2). Strong bias in intercrosses has been reported in previous studies (Table S1). The somewhat variable magnitude of bias between UH and MMCRI may be explained by the differences in mouse source (mice derived recently from frozen embryos at UH versus a longstanding colony at MMCRI; see Materials and Methods), the possibility of microbiome differences in the reproductive tracts between sites22–34, undetected differences in local husbandry conditions, or sampling fluctuations. One or more of these mechanisms may also be responsible for the modest departure from expectations in MMCRI backcrosses, which was evident in both

### Table 1. In vitro fertilization (IVF) and embryo culture for intercrosses and backcrosses. For embryo development after IVF, the results for the reciprocal backcrosses were pooled. Percentage calculations: 2-cell embryos from inseminated oocytes; M/EB and B/ExB from 2-cell embryos. M, morulae; EB, early blastocysts; B, blastocysts; ExB, expanded blastocysts.

| Cross       | No. IVFs | No. oocytes inseminated | No. 2-cell embryos (%) | No. M/EB (%) | No. B/ExB (%) |
|-------------|----------|-------------------------|------------------------|--------------|--------------|
| Intercross  | 9        | 805                     | 456 (57)               | 416 (91)     | 318 (70)     |
| Backcross   | 16       | 972                     | 494 (51)               | 444 (90)     | 324 (66)     |

Three hypotheses could explain departures from Mendelian expectations: Embryonic Lethality, Distorted Transmission, and Biased Fertilization18 (Fig. 1B). Normal litter sizes in intercrosses argue against embryo loss and normal segregation in backcrosses argues against sex-specific distorted transmission38 (see Fig. 1 for a description of the various crosses). Absence of mechanisms to compensate for embryo loss supports this argument. Instead, for these “bias genes”, genotype distortion without reduced litter size argues that fertilization can sometimes be genetically biased towards particular combinations of oocyte and sperm18 (Fig. 1B).

The basis for such biased fertilization is unknown. Especially, it is unclear whether bias results from interactions between gamete-specific factors such as cell-surface receptors and ligands20–22 and/or external factors acquired by sperm during their transit in the epididymis23,24 or by oocytes during their maturation in the oviduct25,26; genetic interactions between factors in eggs and sperm during completion of meiosis at fertilization27,28; or from somatic factors that facilitate fertilization in the oviduct29–31. Here, we used in vitro fertilization (IVF) to test for gamete- versus soma-specific control of biased fertilization. We also measured reproductive features to test alternative explanations for bias and mined available transcriptomes to identify targets of the A1CF RNA-binding protein that could drive haploid selection. We demonstrated that the fertilization bias specifically depends on the genetics of oocyte and sperm and propose that it originates from the functional differences between spermatids arising from sequestration of A1cf transcripts. Previously, evidence for bias was incidental to the primary focus of reported work. Here, we report the first study focused exclusively on the evidence for bias and tests for one of the proposed mechanisms.
male and female backcrosses and thus resulted from alterations of both sperm and oocytes. An attractive possibility is a difference in diets between the two locations—an absence of soybean meal in the UH diet. Soybeans are a significant source of phytoestrogens that could modulate gametogenesis and fertility\cite{35}, possibly serving as an environmental factor promoting biased A1cf segregation.

In the current study no A1cf<sup>mm</sup> homozygotes were obtained in any of the mating crosses on the 129 genetic background. A1cf<sup>mm</sup> homozygotes were observed before among embryos on mixed the 129-B6 genetic background\cite{19} and among offspring on the pure B6 genetic background with a mutation targeting a different side in the A1cf gene (MGI 1917115, mousephenotype.org) (Table S1). It is possible that B6 background carries at least one genetic variant that normalizes fertilization in A1cf<sup>m+</sup> mice thereby enabling production of A1cf<sup>mm</sup> homozygotes. Such genetic variants are extremely common\cite{36}. It is also possible that type of mutation contributed to contrasting results between our study and that reported by mousephenotype.org.

Analysis of three possible sources of distortion supports that Biased Fertilization is responsible for untypical A1cf transmission bias. We next evaluated the evidence for the three possible sources of bias: Embryonic Lethality, Distorted Transmission, and Biased Fertilization (Fig. 1B).

**Embryonic lethality:** Embryonic lethality does not readily account for m+ excess, absence of mm homozygotes, and deficiency of ++ among intercross offspring. Loss of mm should reduce litter size by 25% in intercrosses versus backcrosses where mm homozygotes do not occur. Litter size could be further reduced if some ++ or m+ mice are also lost (Fig. 1B). But in our study litter size did not differ substantially between intercrosses and backcrosses (Fig. 2). At MMCRI, the average litter size for intercrosses and combined backcrosses was 4.9 and 4.6 pups per litter, respectively, whereas at UH these were 6.6 and 6.0, respectively. By comparison, wild-type 129S1/SvImJ inbred mice have an average litter size of 4.9\cite{37}. Finally, an average intercross litter sizes of ~ 8.4–10.0

---

**Figure 2.** A1cf segregation in in vivo test and control crosses. A1cf genotype segregation is shown for two study locations: the Maine Medical Center Research Institute (MMMRI, panel A) and the University of Hawaii (UH, panel B). ++ refers to the wild-type genotype and m+ to the heterozygous genotype. Heterozygotes were not found in either intercross. c² goodness-of-fit tests are shown with point-wise p-values; results remain significant after correcting for testing multiple hypotheses. 'Comment' refers to test results for the fit between total observed genotype numbers with Mendelian expectations for wild-type and heterozygous mice. Sex ratios did not differ significantly from 1:1 expectation (not shown). SEM, standard error of the mean; n, total number of litters.

---

**Table A.**

| Offspring | MMRCRI | Intercross (expect 2 m+ : 1 ++) | Offspring | UH | Intercross (expect 2 m+ : 1 +) |
|-----------|--------|--------------------------------|-----------|----|--------------------------------|
|           |        | WT (++) | Hets (m +) | Ratio | Females | WT (++) | Hets (m +) | Ratio | Females | WT (++) | Hets (m +) | Ratio |
| Females   | 22     | 154     |            | 6.1   | 30     | 75     |            | 3.1   | 40     | 50     |            |       |
| Males     | 29     | 156     |            |       | 22     | 84     |            |       | 41     | 44     |            |       |
| Total     | 51     | 310     |            |       | 52     | 159    |            |       | 81     | 94     |            |       |
| Ave. litter size | 4.9 (SEM=0.30, n=74) | | Ave. litter size | 6.6 (SEM=0.039, n=32) |
| Comment: | Excess heterozygotes. X²*2 = 59.87, | | Comment: | Excess heterozygotes X²*2 = 7.14, |

| Offspring | Female backcross (expect 1 m+ : 1 ++) | Offspring | Male backcross (expect 1 m+ : 1 ++) |
|-----------|--------------------------------------|-----------|-----------------------------------|
|           | MMRCRI | Backcross | UH | | Backcross |
|           | WT (++) | Hets (m +) | Ratio | Females | WT (++) | Hets (m +) | Ratio | Females | WT (++) | Hets (m +) | Ratio |
| Females   | 31     | 53      | 2.2  | 40    | 50     |            |       | 43     | 62     | 1.6    |       |
| Males     | 28     | 77      |       | 41    | 44     |            |       | 38     | 66     | 1.6    |       |
| Total     | 59     | 130     | 2.2  | 81    | 128    | 1.6      |       | 81     | 128    | 1.6    |       |
| Ave. litter size | 4.3 (SEM=0.65, n=44) | | Ave. litter size | 4.8 (SEM=0.63, n=43) |
| Comment: | Excess heterozygotes. X²*2 = 26.67, | | Comment: | Consistent with expectations |

---

**Table B.**
at conception, which is roughly double the observed litter size for backcrosses in the MMCRI study and for the 129S1/SvImJ strain, would be needed to account for the observed loss of all mm and some ++ offspring (Text S2).

Finally, when we evaluated extended MMCRI breeding data, we found that litter sizes were uniformly larger in intercrosses than in backcrosses and the 129S1/SvImJ strain, contrary to the expectation if some mm genotypes were lost postnatally (Table S2). From more than 1300 pups followed between birth and weaning, only six were lost, and all were in backcrosses rather than intercrosses. These lost pups occurred in each of the three types of backcrosses, suggesting loss was random rather than associated with a particular strain combination (Table S2).

Together, these data show that it is unlikely that embryonic lethality contributes significantly to excess heterozygosity.

Ovulation rate provides complementary evidence about embryo loss. The number of live-born offspring is closely related to the number of ovulated oocytes, which is largely determined by female genetics and is typically independent of the genetics of the mating partner. Corpora lutea form where oocytes emerge from ovarian follicles at ovulation and serve as a direct measure of the number of ovulated oocytes. At MMCRI, five intercross females examined at parturition had a total of 41 corpora lutea and 39 embryos. Resorbed intercross embryos were rare, only 1 of 101 embryos across 18 intercross litters, and were not found among embryos flushed from the oviduct at embryonic day 3.5. Finally, in the UH study, none of the reproductive features, including the number of corpora lutea, differed significantly between mm and ++ (Fig. 3). Thus, in neither study location did differences in corpora lutea numbers, resorptions, or other reproductive features account for differences in genotype numbers.

**Distorted transmission.** Transmission Ratio Distortion (TRD) is the preferential transmission of particular alleles from heterozygotes to offspring. These exceptions to Mendelian inheritance are usually sex-specific and independent of the genetics of mating partners. If transmission distortion accounted for bias, similarly strong deviations from Mendelian expectations should be found in backcrosses as well as intercrosses (Fig. 1B).

Distortion observed in some backcrosses (Fig. 2) was too weak to account for the strong bias found in the intercrosses (Text S1).

**Biased fertilization.** For fertilization to cause bias, genotype distortion in embryos or offspring should not be linked to changes in litter size or genotype loss (Fig. 1B). All the data presented above (Table 1, Figs. 2, 3, Text S1–S2, Table S2) agree with these requirements.

Together, the data support that among the three possible sources of distortion, the evidence is consistent only with Biased Fertilization.
Transcriptome data mining points to a possible mechanism for fertilization bias in A1cf transmission. What factors intrinsic to sperm might predispose them to bias? A1cf and four of the nine currently known “bias genes” encode RNA-binding proteins (RBPs) that control many aspects of RNA biology (Fig. 4A). A1CF is one of the mRNA targeting components of the APOBEC1 RNA editing complex40 and also controls splicing41. A1CF protein expression is high in mature oocytes at all ages examined as well as in testis, spermatozoa and epididymides, but declines with age in testes and epididymides42. AGO2 is part of the RISC complex that silences miRNAs, piRNAs and transposons43. DDX1 is an ATP-dependent RNA DEAD-box helicase involved in translation, splicing, and miRNA processing44. DND1 blocks miRNAs access to their mRNA targets thereby stabilizing target transcripts for translation45. PUM1 represses translation by binding 3′ UTR’s of target transcripts, promoting miRNA access, and modulating mRNA stability46. The question is whether localizations of these RBPs or their targets is restricted to the gametes in which they originate, thereby creating opportunities for functional variation that could be the basis for haploid selection and biased fertilization.

To address this question, we examined previously published transcriptomes data12 for localized mRNA expression in spermatids from wild-type mice. This survey found more than 12,000 genes, including ‘genoinformative markers’ (GIMs) whose RNAs are enriched in the spermatid where they were transcribed as well as non-GIMs whose RNAs shuttle among connected spermatids. Interestingly, four of the eight “bias genes” are GIMs (A1cf, Lrp6, Pum1, Vangl2), two are non-GIMs (Ddx1, Dnd1), and two are ‘remaining genes’ (gray). Right: the kernel density showing that most mouse genes had low genoinformativity scores.

Figure 4. RNA-binding proteins and genoinformativity markers. (A) RNA-binding proteins (RBPs) and genoinformativity markers (GIMs). Source of information: Chromosome (informatics.jax.org), RBPs18, and GIMs, non-GIMs and sweeps12. The approximate distance (kb) of “bias” genes from the center of the sweep is provided in parentheses. The Ppp2cb transcript was detected but did not reach the threshold to determine whether it is a GIM or non-GIM12. The ninth ‘bias’ gene, apolipoprotein B transcript (ApoB), which is not known to be an RBP, was not detected12. (B) Genoinformativity scores. Left: genoinformativity scores and 95% confidence intervals12 are displayed. Four are confident GIMs (red), two confident non-GIMs (black), and two “remaining genes” (gray). Right: the kernel density showing that most mouse genes had low genoinformativity scores.
genes in mice with a liver-specific A1cf deficiency \(^41\) (\(\chi^2 = 6.40, 2\) df, \(p < 0.05\); Fig. S1). Similar analyses for enriched binding motifs and functions among mRNA targets of the other RBP GIM (Pum1) did not reveal notable trends (Fig. S2). Finally, several genes that interact directly with "bias genes" or that show altered function in mice with a mutation in a "bias gene" show non-Mendelian segregation (Table S5, Fig. S3). Collectively, these genes define a network of protein interactions and functional dependencies that coordinate mRNA stability, content, and translation control (Fig. 5). We propose that loss of sequestered transcripts in spermatids carrying the A1cf loss-of-function allele leads to anomalous transcripts and protein products, and in turn to functional variation in both m and + spermatids in m+ heterozygotes. Similar effects might be found in mature m+ oocytes where A1cf is highly expressed \(^42\).

Because variants for the nine "bias genes" are either engineered or spontaneous mutations \(^18\), they have not been subject to natural selection. One could therefore argue that disrupting fundamental molecular features informs the mechanisms of fertilization in laboratory settings but does not necessarily guide understanding about the genetics of natural populations. We therefore asked whether "bias genes" have experienced positive selection, as expected for genetic variants that distort transmission. Several GIMs in mice and primates show evidence of selective sweeps in humans \(^12\), where the frequency of the selected genetic variant increases because of its fitness advantage \(^47\). Closely linked genetic variants hitchhike with the selected locus, showing correlated gene frequency changes depending on their recombination distance from the selected locus, and leaving a signature of reduced genetic variation. Interestingly, human orthologues of three "bias genes" (Ago2, Dnd1, Ppp2cb) are significantly enriched in selective sweeps (\(p < 0.0004\); Fig. 4A), but whether "bias genes" drive these sweeps is uncertain.

**Discussion**

Chance is central to sexual reproduction, from segregation and recombination to the union of gametes at fertilization. Although precise molecular mechanisms guide meiosis, recombination, and fertilization \(^48\)–\(^50\), the unique combination of crossovers on each chromosome, chromosomes allocated to each gamete, and gametes that join at fertilization together make the genetic outcomes seemingly unpredictable. The statistical properties of Mendel’s Laws emerge from this mix of precision and chance. Exceptions such as transmission distortion drive preferential occurrence of alleles to the next generation, thereby reducing genetic variation and making outcomes more predictable.

**Genetics of gametes versus gene-environment interaction.** A key question is whether the determinants of bias are intrinsic to gametes, or instead depend on gene-environment interactions. Factors in semen and from uterus, oviduct and ovarian follicles promote gamete viability, enhance chemotaxis and thermotaxis, and facilitate fertilization \(^29\)–\(^31\),\(^51\)–\(^52\). However, if these factors alone were sufficient for bias, gametes would be similarly affected regardless of their genetic content. Experimental in vivo approaches to this question face formidable challenges including the limited number of mature oocytes and sperm at fertilization, the transient nature of active gametes, and the ephemeral environment of the oviduct at fertilization. IVF obviates many of these challenges. The results reported here clearly show that bias depends on gamete genetics alone, or perhaps on a genetically-determined predisposition to environmental change that arises earlier during gametogenesis and persists after gametes are isolated for IVF tests.
Functional consequences of bias genes. The function of bias genes can provide insights into molecular and cellular mechanisms that lead to the non-random union of gametes. During gametogenesis, some RNAs and proteins tend to remain within the spermatid in which they originate rather than translocate through cytoplasmic bridges that connect the four gametes resulting from each meiosis. RBPs shuttle mRNAs to germ granules, such as chromatoid bodies, where their cargos can be stored, processed, or transported intra-cellularly and sometimes across these bridges. Sequestered transcripts and proteins sometimes create functional differences among genetically distinct but otherwise phenotypically indistinguishable spermatids. Any resulting phenotypic differences can be the basis for competition among gametes that sometimes leads to transmission distortion.

Interestingly, "bias genes" appear to be connected in a network of functional dependencies and protein interactions that coordinates mRNA stability, editing, and translation control with various germline processes (Fig. 5).

Model. Any model for biased fertilization must accommodate several observations. Sperm and oocytes carrying intrinsic functional differences in A1cf heterozygotes arrive at the site of fertilization in appropriate numbers and functionality, as normal segregation in the in vivo and in vitro backcrosses demonstrate. Most ovulated oocytes are fertilized, leaving little opportunity for selection among oocytes. Finally, the window for bias in females is remarkably short, between the time when sperm trigger resumption of the second meiotic division (MII) and fusion of the female and male pronuclei. Two models could account for biased fertilization (Fig. 6). The first involves Oocyte Selection where sperm bearing the A1cf+ allele selectively fertilize mm oocytes while those bearing the m allele fertilize ++ oocytes. Perhaps isoform variants within a species confer specificity to oocyte-sperm interactions as they do between species. Alternatively, bias could result from a reversed order of meiotic divisions during oogenesis thereby creating an opportunity to direct chromatid segregation at MII and whereby retain a preferred chromatid in the oocyte, depending on the genetics of fertilizing sperm (Fig. 6, Reverse Meiosis). With the usual order of meiotic divisions where the reductional division precedes the equational division, preferential retention of recombinant chromosomes could augment or reduce the magnitude of bias. Several genes that regulate meiosis were found among A1cf targets in spermatids. If enrichment is genotype-specific, these transcripts might influence a completion of female meiosis post-fertilization. Reversed (inverted) meiosis has been reported in humans and mice and plant species with holocentric chromosomes. In both models, preferential retention of recombinant chromosomes could augment or reduce the magnitude of bias. Tests for these models are underway.
Implications. In often elaborate ways diploid organisms choose mates; perhaps haploid gametes do the same. As Darwin proposed, mate choice depends on the perceived fitness of potential partners—fit mates tend to produce fit offspring. But gametes in such individuals are not equally fit. Although precise molecular mechanisms guide meiosis, recombination and fertilization, chance determines the combination of crossovers on each chromosome and chromosomes allocated to oocytes and sperm, making each gamete genetically unique. Sequestration of mRNAs and their protein products creates further functional diversity. As a result, fitness can vary among gametes even in fit individuals. Non-random joining of oocytes and sperm at fertilization could reflect haploid selection for fit combinations of gametes. Future in vitro as well as in vivo studies with A1cf and the other bias genes will help to clarify this interplay between chance and selection in gamete preference at fertilization.

Materials and methods

Mice. Work at both institutions used the same mutation (129/Sv-A1cf<sup>tm1Ddsn</sup>/NaJ) that was made and maintained on the 129S1/SvImJ inbred background (Jackson Laboratory, Strain #: 002440<sup>19,42</sup>). Mice for work at MMCRI were obtained from JNH's breeder colony. Some of these mice were also donated to the Jackson Laboratory as a resource for the research community. Frozen embryos from the Jackson Laboratory were used to establish the UH breeder colony.

The A1cf (A1cf<sup>tm1Ddsn</sup>, MGI:3690371) mutants were made with a strain 129-derived ES cell line<sup>19</sup> and then rederived on the 129S1/SvImJ inbred background, without outcrossing with C57BL6J or other strains<sup>42</sup>. Isogenicity greatly reduces the possibility of extraneous genetic effects. These mutant mice were provided to the Jackson Laboratory for preservation and distribution (Jackson Laboratory, Strain #: 027924). Mice at MMCRI were maintained on Envigo Teklad 2918 irradiated Chow diet (Envigo RMS, Indianapolis, IN) provided ad libitum, at 70°F and with a 14L:10D schedule, with light-on at 7 am and light-off at 9 pm, Alpha Dry Plus bedding (Shepherd Specialty Paper, Watertown, TN), and in static cages (Allentown Inc., Allentown, NJ).

At UH, frozen embryos (A1cf<sup>tm1Ddsn</sup>, MGI:3690371) were acquired as a mixture of wild-type and heterozygous embryos (Jackson Laboratory, Strain #: 027924). Embryos were transferred into oviducts of CD-1 pseudopregnant recipients and resided in heterozygous males and females that were used to establish a breeder colony. Mice were fed <i>ad libitum</i> with a standard diet (2020X Teklad Global Soy Protein-Free Extrудed Rodent Diet, Harlan Laboratories, Indianapolis, IN) and were maintained with 7092 Teklad Corn cob bedding (Envigo, Indianapolis, IN) in a temperature- and light-controlled room (22 °C, 14L:10D, with light-on at 7 am and light-off at 9 pm) in accordance with the guidelines of the Laboratory Animal Services at the University of Hawaii and guidelines presented in National Research Council’s Guide for Care and Use of Laboratory Animals published in 1996 by Institute for Laboratory Animal Research (ILAR) of the National Academy of Science (Bethesda, MD).

Experimental protocols for animal work were approved by the Maine Medical Center Research Institute Institutional Animal Care and Use Committee (protocols 1911 and 1912) or by the University of Hawaii Institutional Animal Care and Use Committee (protocol 06-010). The reporting in the manuscript follows the recommendations in the ARRIVE guidelines.<sup>58</sup>

Media. Mineral oil was purchased from FUJIFILM Irvine Scientific (Santa Ana, CA); pregnant mares’ serum gonadotrophin (eCG) and human chorionic gonadotrophin (hCG) from ProSpec (East Brunswick, NJ). All other chemicals were obtained from Sigma Chemical Co. (St Louis, MO) unless otherwise stated. Medium HTF was used for IVF<sup>69</sup>, Heparin-buffered CZB (HEPES-CZB) for gamete handling<sup>66</sup>, and mKSOM<sup>AA</sup> (KSOM medium containing NEAA and EAA<sup>62</sup> for embryo culture. Both mKSOM<sup>AA</sup> and HTF were maintained in an atmosphere of 5% CO<sub>2</sub> in air and HEPES-CZB was maintained in air.

In vitro fertilization and embryo culture. Sperm capacitation and IVF were performed as previously reported<sup>69</sup>. Briefly, oocytes were collected from females induced to superovulate with injections of 5 IU eCG and 5 IU hCG given 48 h apart. Epididymal sperm were collected by release from cauda epididymis directly into HTF medium and were capacitated for 1.5 h at 37 °C in a humidified atmosphere of 5% CO<sub>2</sub>. Gametes were co-incubated for 4 h, after which oocytes were washed with HEPES-CZB, followed by at least one wash with mKSOM<sup>AA</sup> medium. Morphologically normal oocytes were selected for culture. Fertilized oocytes (oocytes with two well-developed pronuclei and extruded 2nd polar body) were cultured in 50 μL drops of medium containing NEAA and EAA<sup>62</sup> for embryo culture. Both mKSOM<sup>AA</sup> and HTF were maintained in an atmosphere of 5% CO<sub>2</sub> in air and HEPES-CZB was maintained in air.

Mouse genotyping. MMCRI. PCR was used to amplify A1cf alleles as recommended by the Jackson Laboratory for some mice or with an in-house protocol for others. Genomic DNA was extracted from toe clippings with this use of the Hot Shot protocol (L. Liaw Lab, MMCRI; https://health.uconn.edu/mouse-genome-modification/protocols/hotshot-method-of-dna-preparation). Samples were exposed to a lysis buffer (25 mM NaOH, 0.2 mM EDTA, pH 12) at 95 °C for 45 min. before the extract was neutralized with 40 mM Tris HCl, pH 5.

Two PCR protocols were used: the Jackson Laboratory protocol and the in-house amplification protocol. The Jackson Laboratory protocol used three primers: a wild-type forward primer (24,997: GTC CAG GGC ACA GGA TAT AG), a mutant forward primer (oIMR7415: GCC AGA GGC CAC TTG TGT AG), and a common reverse primer (24,998: CAA AGG TCT AAC ATG GGA TAT AG), a mutant forward primer (oIMR7415: GCC AGA GGC CAC TTG TGT AG), and a common reverse primer (24,998: CAA AGG TCT AAC ATG GGA TAT AG). Amplifications were performed with positive, negative, and blank controls. Because all samples were used to yield a product, amplification control was not included. All reactions were performed in 30 μL, with overall primer concentrations of 0.33 μM for 24,997 and 24,998, and 0.40 μM for oIMR7415 in Taq Red Master Mix Kit (1.5 mM MgCl<sub>2</sub>) (Genesee Scientific, El Cajon, CA).
The experiments were designed to compare observed and expected genotype distributions for intercrosses and backcrosses (Fig. 1A) as well as for embryos after IVF.

In the breeding studies, the following outcomes were measured: (1) litter size; (2) m:++ ratio; and (3) sex ratio (the numbers of females and males). In the IVF study, the following outcomes were measured: (1) number of oocytes per oviduct; (2) the proportion of 2-cell embryos obtained from oocytes inseminated; (3) the proportion of morulae and blastocysts obtained from cultured 2-cell embryos; (4) the blastocyst m:++ ratio; and (5) homozygote viability and genotyping results. The experiments were designed to compare observed and expected genotype distributions for intercrosses and backcrosses (Fig. 1A) as well as for embryos after IVF.
the blastocyst sex ratio. The additional measures included analyses of m+ and ++ A1cf females (body weight, ovary weight, corpora lutea scores) and m+ and ++ A1cf males (body weight, testis weight, sperm number, sperm motility).

Depending on the study, chi-square goodness-of-fit or chi-square contingency tests were used, both with 1 degree of freedom. Statistical significance was claimed for tests showing \( p < 0.05 \), both point-wise as well as after experiment-wide Bonferroni correction for multiple testing. The test score and its point-wise probability are provided; the results remain significant after accounting for testing six crosses. Data analysis and computations were done with Excel and GraphPad Prism.

**Network analysis.** Several databases were queried to identify candidate partners of that interact directly with “bias proteins” (protein–protein interactions), that show functional dependency in mutant mice, or that are no more than two nodes away from “bias proteins” in interaction networks. The evidence was derived from the primary literature or from various genomic databases (uniprot.org, string-db.org). Given the incomplete nature of the surveys for interactions, and the lack of systematic studies across tissues, stages of development, environmental conditions, and genetic backgrounds, it is possible that some interactions have not yet been documented.

In some cases, at least one member of a gene family was counted as a putative interactor with a bias protein. For example, DDX1 interacts with PPP1R8, but the literature is silent about interactions with PPP2CB (string-db.org). Because PPP2CB contributes to control of meiosis in oogenesis, we connect DDX1 with the PP2 family and mark the interaction as ‘provisional’. Similarly, A1CF interacts with at least one member of the elf4 translation family but interactions have not been tested with other members of the family. Finally, only those interactions were included that provide evidence for interactions among the bias genes that nucleate the network.

Segregation data were obtained from mousephenotype.org or the literature.

**Motif enrichment.** For each gene expressed (TPM > 10) in mouse spermatids in Bhutani et al., we retrieved the annotated 3’ UTR sequences for the highest-expressed Ensembl transcript. Using canonical iUPAC motifs (WUAUUR for A1cf and UGUANAU for Pum1), we searched for enrichment using AME from the MEME Suite. Position weight matrix (PWM) was used to evaluate the match between candidate RNA targets of A1cf and Pum1 motif sequences. We searched for enrichment in GIMs against the control background of expression-matched spermatid-expressed controls (all 20 sets) as reported in Bhutani et al. As an additional control, we searched for enrichment in the first control set against the other 19 control sets and verified that the adjusted \( p \)-values for each motif were greater than 1. To classify genes as GIM’s, non-GIM’s or remaining genes, we selected cutoffs on mean genoinformativity, confidence in the genoinformativity estimate, and confidence in haplotype calling for that region of the chromosome, that would in aggregate lead to a false discovery rate of no more than 10%. See Bhutani et al. for details.

**GO ontology enrichment analysis.** Enrichment analysis of binding partners for A1cf and Pum1 for gene ontology (GO) terms was performed using the goana function in the limma package using all genes as the denominator. Results were truncated at \( p < 0.01 \) for A1cf (\( n = 121 \)) and \( p < 0.001 \) for Pum1 (\( n = 56 \)). The threshold was more stringent for Pum1 than A1cf because the collective results were highly heterogeneous, with few themes beyond ‘transcription’. Descriptors for these prioritized GO terms were distilled into several higher-level summary terms, e.g. ‘RNA included functions such as splicing, RNA transport and polyadenylation, and ‘meiosis’ included recombination and centrosomes. ‘Other’ included heterogeneous functions. The GO term id’s and descriptors were updated with the corresponding GO hierarchies and gene lists (Mouse Genome Database, informatics.jax.org), especially in cases where the descriptor was general, e.g. metabolic process (GO:0008152). In some cases, the primary literature was also reviewed.

**Selective sweep.** Data were obtained from Bhutani et al. To calculate the probability of finding three of nine “bias genes” in selective sweeps, we assumed that genome-wide 100 sweeps involved a total of 50 MB (average 500 kb per sweep) corresponding to a probability of 0.017 that any three “bias genes” occur in the total area.

**Data availability**

All data are available in the main text or the supplementary materials.

Received: 11 May 2022; Accepted: 3 August 2022
Published online: 10 August 2022

**References**

1. Bateson, W. & Mendel, G. Mendel’s Principles of Heredity (Cambridge University Press, 1902).
2. Crow, J. F. Why is Mendelian segregation so exact? BioEssays 13, 365–312. https://doi.org/10.1002/bies.950130609 (1991).
3. Agren, J. A. & Clark, A. G. Selfish genetic elements. PLoS Genet 14, e1007700. https://doi.org/10.1371/journal.pgen.1007700 (2018).
4. Immler, S. & Otto, S. P. The evolutionary consequences of selection at the haploid gametic stage. Am. Nat. 192, 241–249. https://doi.org/10.1086/698483 (2018).
5. Braun, R. E., Behringer, R. R., Peschon, J. J., Brinster, R. L. & Palmer, R. D. Genetically haploid spermatids are phenotypically diploid. Nature 337, 373–376. https://doi.org/10.1038/337373a0 (1989).
6. Erickson, R. P. Haplotype gene expression versus meiotic drive: the relevance of intercellular bridges during spermatogenesis. Nat. New Biol. 243, 210–212. https://doi.org/10.1038/newbio243210a0 (1973).
7. Soygur, B. et al. Intercellular bridges coordinate the transition from pluripotency to meiosis in mouse fetal oocytes. Sci. Adv. https://doi.org/10.1126/sciadvabc6747 (2021).
50. Kornbluth, S. & Fissore, R. Vertebrate reproduction. *Cold Spring Harb. Perspect. Biol.*, 7, a006064. https://doi.org/10.1101/cshperspect.a006064 (2015).
51. Eisenbach, M. & Giojalas, L. C. Sperm guidance in mammals—an unpaved road to the egg. *Nat. Rev. Mol. Cell Biol.*, 7, 276–285. https://doi.org/10.1038/nrm1893 (2006).
52. Kekalainen, J. & Evans, J. P. Female-induced remote regulation of sperm physiology may provide opportunities for gamete-level mate choice. *Evolution*, 71, 238–248. https://doi.org/10.1111/evol.13141 (2017).
53. Kotaja, N. & Sassone-Corsi, P. The chromatoid body: a germ-cell-specific RNA-processing centre. *Nat. Rev. Mol. Cell Biol.*, 8, 85–90. https://doi.org/10.1038/nrm2081 (2007).
54. Haldane, J. A mathematical theory of natural and artificial selection. Part II. The influence of partial self-fertilisation, inbreeding, and selective fertilisation of Mendelian populations, and natural selection. *Biol. Rev.*, 1, 158–163 (1924).
55. Gert, K. R. & Pauli, A. Species-specific mechanisms during fertilization. *Curr. Top. Dev. Biol.*, 140, 121–144. https://doi.org/10.1016/bs.ctdb.2019.10.003 (2020).
56. Cabral, G., Marques, A., Schubert, V., Pedroza-Harand, A. & Schlegelhofer, P. Chiasmatic and achiasmatic inverted meiosis of plants with holocentric chromosomes. *Nat. Commun.*, 5, 5070. https://doi.org/10.1038/ncomms6070 (2014).
57. Singh, N. D. *et al.* Fruit flies diversify their offspring in response to parasite infection. *Science*, 349, 747–750. https://doi.org/10.1126/science.aab1768 (2015).
58. Kilkenny, C., Browne, W. J. I., Cuthill, I., Emerson, M. & Altman, D. G. Improving bioscience research reporting: the ARRIVE guidelines for reporting animal research. *Vet. Clin. Pathol.*, 41, 27–31. https://doi.org/10.1111/j.1939-165X.2012.00418.x (2012).
59. Quinn, P. Enhanced results in mouse and human embryo culture using a modified human tubal fluid medium lacking glucose and phosphate. *J. Assist. Reprod. Genet.*, 12, 97–105. https://doi.org/10.1007/BF02221377 (1995).
60. Chatot, C. L., Zionek, C. A., Bavister, B. D., Lewis, J. L. & Torres, I. An improved culture medium supports development of random-bred 1-cell mouse embryos in vitro. *J. Reprod. Fertil.*, 86, 679–688. https://doi.org/10.1530/jrf.0.0860679 (1989).
61. Quinn, P., Barros, C. & Whittingham, D. G. Preservation of hamster oocytes to assay the fertilizing capacity of human spermatozoa. *J. Reprod. Fertil.*, 66, 161–168. https://doi.org/10.1530/jrf.0.0660161 (1982).
62. Summers, M. C., McGinnis, L. K., Lawitts, J. A., Raffin, M. & Biggers, J. D. IVF of mouse ova in a simplex optimized medium supplemented with amino acids. *Hum. Reprod.*, 15, 1791–1801. https://doi.org/10.1093/humrep/15.8.1791 (2000).
63. Ajduk, A., Yamauchi, Y. & Ward, M. A. Sperm chromatin remodeling after intracytoplasmic sperm injection differs from that of in vitro fertilization. *Biol. Reprod.*, 75, 442–451. https://doi.org/10.1095/biolreprod.106.053223 (2006).
64. Su, Y. Q. *et al.* MARF1 regulates essential oogenic processes in mice. *Science*, 335, 1496–1499. https://doi.org/10.1126/science.1214680 (2012).
65. McLeay, R. C. & Bailey, T. L. Motif Enrichment Analysis: a unified framework and an evaluation on ChIP data. *BMC Bioinform.*, 11, 165. https://doi.org/10.1186/1471-2105-11-165 (2010).
66. Castle, W. E. & Little, C. C. On a modified mendelian ratio among yellow mice. *J. Reprod. Fertil.*, 32, 833–868 (1910).

**Acknowledgements**

We thank Valerie Blanc, Nicholas Davidson, Kari Koszdin, Alma Rivera and Jacqui White for sharing information about the diet on which mice were bred; Jason Heaney for providing information about the nature of the A1cf mutant that IMPC produced; and Kaitlin Nee for maintaining the mouse colonies and Chloe Kumpel for assisting with genotyping and data management at MMCRI. The Histology Core at the University of Hawaii was supported by National Institutes of Health (NIH) Grant U54MD007601.

**Author contributions**

Conceptualization: J.H.N., M.A.W.; Methodology: J.H.N., M.A.W., R.F.; Formal analysis: J.H.N., M.A.W., R.F., C.Y.K., C.W.L.; Investigation: N.H., G.B., Y.Y., A.S.; Resources: J.H.N., M.A.W.; Data curation: J.H.N., M.A.W., R.F.; Writing–original draft: J.H.N.; Writing–review and editing: J.H.N., M.A.W., N.H., G.B., Y.Y., R.F., C.Y.K., C.W.L., A.S.; Funding acquisition: J.H.N., M.A.W.; Project administration: J.H.N., M.A.W.

**Funding**

National Institutes of Health grant R56HD098217 (JHN). National Institutes of Health R01HD072380 and R03HD106936, and Hawaii Community Foundation 20CON-106807 Grants (MAW).

**Competing interests**

The authors declare no competing interests.

**Additional information**

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1038/s41598-022-17948-9.

**Correspondence** and requests for materials should be addressed to M.A.W. or J.H.N.

**Reprints and permissions information** is available at www.nature.com/reprints.

**Publisher’s note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
