Workflow Optimization for Identification of Female Germline or Oogonial Stem Cells in Human Ovarian Cortex Using Single-Cell RNA Sequence Analysis

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Abstract

In 2004, the identification of female germline or oogonial stem cells (OSCs) that can support post-natal oogenesis in ovaries of adult mice sparked a major paradigm shift in reproductive biology. Although these findings have been independently verified, and further extended to include identification of OSCs in adult ovaries of many species ranging from pigs and cows to non-human primates and humans, a recent study rooted in single-cell RNA sequence analysis (scRNA-seq) of adult human ovarian cortical tissue claimed that OSCs do not exist, and that other groups working with OSCs following isolation by magnetic-assisted or fluorescence-activated cell sorting have mistaken perivascular cells (PVCs) for germ cells. Here we report that rare germ lineage cells with a gene expression profile matched to OSCs but distinct from that of other cells, including oocytes and PVCs, can be identified in adult human ovarian cortical tissue by scRNA-seq after optimization of analytical workflow parameters. Deeper cell-by-cell expression profiling also uncovered evidence of germ cells undergoing meiosis-I in adult human ovaries. Lastly, we show that, if not properly controlled for, PVCs can be inadvertently isolated during flow cytometry protocols designed to sort OSCs because of inherently high cellular autofluorescence. However, human PVCs and human germ cells segregate into distinct clusters following scRNA-seq due to non-overlapping gene expression profiles, which would preclude the mistaken identification and use of PVCs as OSCs during functional characterization studies.

Key words: human ovary; single-cell RNA sequence analysis; oocyte; germ cell; oogonial stem cell.
Introduction

A central underpinning of reproductive biology has held that oocyte generation in ovaries of female mammals is restricted to the embryonic period. This thinking deviates markedly from spermatogenesis in males throughout adult life, which involves meiotic differentiation of male germline or sperm-agonial stem cells (SSCs) in the testes. However, the longstanding paradigm of a non–renewing oocyte pool was challenged by a study with mice in 2004, which reported the existence of female germline or oogonial stem cells (OSCs) and the continuation of oocyte production in adult mouse ovaries. While this study sparked significant debate, more than 80 corroborating studies now support the existence of OSCs and/or active oogenesis in adulthood across species, including humans (Supplementary Tables S1–S3). The discovery of OSCs, which brings the biology of male and female gametogenesis in mammals more closely in line with one another and with that of non-vertebrate species, has significant ramifications for the development of in vitro models to investigate human oogenesis as well as for new technologies to combat ovarian failure and female infertility caused by aging or insults.

A major breakthrough in the study of OSCs came in 2009, with the first report that the cells could be retrieved as a distinct population from mouse ovaries using DEAD-box helicase 4 (DDX4) antibody-based sorting. Through extensive in vitro characterization and in vivo transplantation studies, the germline identity of the cells was established, as was the functional identity of the cells as bona fide precursors to oocytes that can be fertilized to produce viable offspring.

More than 60 other publications have since isolated OSCs from ovaries of mice, rats, pigs, cows, baboons, and humans (Supplementary Tables S1 and S2). Moreover, inducible genetic lineage tracing studies with mice have fate–mapped new oocytes produced during adulthood to the generation of healthy offspring in natural mating trials, thus establishing the physiological importance of OSC-supported oogenesis to adult ovarian function and female fertility.

A second major breakthrough came in 2012 with the successful purification of OSCs from adult human ovarian cortical tissue, the findings of which have since been independently verified and extended by many other groups. Human OSCs express a profile of genes characteristic of primitive germ cells, and these cells differentiate through meiosis into oocyte–like cells in vitro and into oocytes that are enclosed within newly formed follicles after transplantation into human ovarian cortical tissue.

Human OSCs isolated by fluorescence-activated cell sorting (FACS) with monoclonal antibodies against DDX4 have also been used in approved clinical studies. Discordant with this large body of work, a recent study concluded from single–cell RNA sequence analysis (scRNA-seq) that human OSCs do not exist. These authors identified only 6 clusters (viz. 6 cell types) in adult human ovarian cortical tissue biopsies using scRNA-seq: stromal cells, perivascular cells (PVCs), endothelial cells, granulosa cells, immune cells, and oocytes. That non–oocyte germ lineage cells were apparently missing from their analysis was subsequently put forth as evidence of OSCs being absent from adult human ovaries. However, the expression and clustering analysis reported in this study were performed with an early version of Cell Ranger software (2.1.1 or v2), which has widely known limitations in its ability to detect low-expression cells. An improved version of Cell Ranger software (3.0.2 or v3), which was available and actually used by the authors in the same study for analysis of human ovarian cells after flow cytometric sorting, increases the sensitivity for cell calling by approximately one log-order over that using Cell Ranger v2 (https://www.10xgenomics.com). While the preprocessed data obtained from Cell Ranger v2 and v3 are fairly consistent, the ability of Cell Ranger v3 to detect more cells, especially those with low abundance transcripts, can change batch-effect corrections and thus the accuracy of the output data analysis. Another issue that can affect the resolution of scRNA-seq is the human reference genome used for read alignments, with HG38 preferred over HG19 for optimal depth of analysis. All of this is highly relevant because OSCs, like other stem cell types, are very rare, with a reported frequency of ~0.014% in adult ovaries. If one is seeking to identify as many cells, and as many cell types, as possible in a highly heterogenous cell sample using scRNA-seq, decisions about which analytical approaches will be used become critically important to consider prior to performing the experiments.

In this same study, Wagner et al. also used DDX4 antibody-based FACS coupled with scRNA-seq to claim that OSCs isolated and studied by many others for over a decade using the same cell sorting strategy have been critically important to consider prior to performing the experiments.

In parallel, we evaluated the possibility that technology like scRNA-seq could provide further evidence of not just the existence of human OSCs but also of primitive germ cells committing to, or progressing through, the early stages of meiotic differentiation into oocytes in adult human ovarian tissue under normal physiological conditions in vivo. Such an outcome, which has not yet been demonstrated in humans, would be consistent with recent genetic tracing studies in adult female mice showing that active meiotic entry and oogenesis occur naturally in adult ovaries during reproductive life, and that oocytes formed in the ovaries during adulthood contribute directly to the pool of eggs used for natural species propagation.
Materials and Methods

Animals
Freshly collected ovarian tissues from adult heifers (Bos taurus) were obtained from Blood Farms (Groton, MA) and processed immediately for cortex isolation and cryopreservation until use.

Human Subjects
All research with human tissues was approved by the institutional review boards of Northeastern University (IRB#14-03-22), University of Edinburgh (LREC 16/SS/0144), and Saitama Medical University (630-III). Informed consent was obtained from all participants, and all tissue samples were de-identified prior to use. A total of 7 ovarian cortical tissue samples from 2 caesarean section patients (CSP) and 1 gender reassignment patient (GRP) between 26 and 34 years of age were used.

Adult Human Unsorted Ovarian Cortical Cell scRNA-seq Data and Code Availability
The scRNA-seq data referenced in this study were originally generated by Wagner et al.25 from adult human ovarian cortical biopsies of 4 subjects (CSP, n = 3; GRP, n = 1). The 10x Genomics dataset of Wagner et al.25 for adult human unsorted ovarian cortical cells was deposited by these authors to, and accessed by us through, the European Molecular Biology Laboratory’s European Bioinformatics Institute (EMBL-EBI) under the accession code E-MTAB-8381. Analyses of scRNA-seq data were completed using the lines of code for adult human unsorted ovarian cortical cells deposited by Wagner et al.25 on GitHub (https://github.com/wagmag/SingleCellOvary). For Cell Ranger v6 analysis, additional lines of code were run in parallel to select proper quality control metrics as well as to determine the parameters for dimensionality reduction that best represented the data. The code used with the Cell Ranger v6 analysis is available on GitHub (https://github.com/hanrico/Ovarian-scRNA-seq).

Clustering and Analysis of scRNA-seq Data
In the Wagner et al.25 study, output files for their adult human unsorted ovarian cortical cell samples were converted using Cell Ranger v2. We first re-analyzed their raw fastq files using the same version of Cell Ranger and the same human genome assembly (HG19), along with Seurat version 3.0.0 (v3) and the lines of code for unsorted human ovarian cortical cells required to detect a given cell type with scRNA-seq.

Flow Cytometry
Ovarian cortical tissue from adult heifers or reproductive-age women was cryopreserved, thawed, and dissociated into single-cell suspensions for flow cytometry using a BD FACSAria III, as described previously (see Supplementary material, Method 2 for more details). Primary antibodies against SMA (ab5694, 1:50; Abcam, Cambridge, MA) or CD31 (MA3100, 1:50; Invitrogen–ThermoFisher Scientific, Waltham, MA), each directly conjugated to APC (Abcam, ab201807), were used for detection of the total percentage of autofluorescent events that were positive for expression of either PVC marker. Flow cytometry data were acquired using BD FACSDiva 8.0.1 software and analyzed by FlowJo 10.7 software.

Data Analysis
For scRNA-seq data, analysis was performed using the human unsorted ovarian cortical cell dataset of Wagner et al.25 For FACS, the data represent the mean ± SE of combined results; n = 4 (CD31) or 7 (SMA), and n = 5 (SMA) or 6 (CD31), for bovine and human ovarian tissue sample analysis, respectively.

Results
Analysis of Unsorted Cells Isolated from Human Ovarian Cortical Tissue Biopsies
We first used the reported frequency of OSCs in adult ovarian tissue with software for estimating the number of cells required to detect a given cell type with scRNA-seq (www.satijalab.org/howmanycells). Assuming 6 general cell types based on the clusters reported by Wagner et al.25 we determined that 84 550 viable ovarian cells would be needed for detection of at least 5 OSCs at 95% confidence (Supplementary Fig. S1). Any number less than 5 cells failed to produce a reliable assessment of input cell number required. Notably, the 12 160 cells analyzed by Wagner et al.25 was 14.4% of the minimal cell input number needed for detection.
Analysis of Human Unsorted Ovarian Cortical Cells for Evidence of Oocytes

Using our optimized analytical pipeline, 10 cells were identified in the germ cell/oocyte cluster as oocytes based on co-expression $\text{FIGLA}$, $\text{OOSP2}$, $\text{GDF9}$, and $\text{ZP3}$ in each cell (Fig. 1F). While $\text{FIGLA}$ is often referred to as an oocyte-specific gene, $\text{GDF9}$ encodes a transcription factor that functions at various stages of oogenesis, including the regulation of key genes needed for meiosis-I progression in premeiotic germ cells. We, therefore, performed a gene-by-gene analysis of the 62 cells in this cluster using Loupe Browser. We identified a total of 40 $\text{FIGLA}$-expressing cells, of which only 21 co-expressed $\text{OOSP2}$, 13 co-expressed $\text{GDF9}$, 37 co-expressed $\text{ZP3}$, and 12 co-expressed both $\text{GDF9}$ and $\text{ZP3}$ (Fig. 2). Likewise, when the other oocyte markers were analyzed individually, we identified 25 $\text{OOSP2}$-expressing cells, 14 $\text{GDF9}$-expressing cells, and 42 $\text{ZP3}$-expressing cells in this cluster (Fig. 2). The discordance in numbers of cells expressing each gene individually or in pairs versus combined as a 4-gene panel may be due to differences in the timing of expression of the various genes relative to oocyte maturational stage, expression in cells other than oocytes, and/or degradation of mRNA transcripts during sample processing that led to the expression of a given gene in a given cell falling below the detection threshold (see Discussion and Supplementary material, Discussion 1 for additional details).

While $\text{DDX4}$ is widely accepted as being expressed in all oocytes in vivo, only 12 of the 62 cells in this cluster expressed $\text{DDX4}$ (Fig. 3). Moreover, only 5 of the $\text{FIGLA}$/$\text{OOSP2}$/$\text{GDF9}$/$\text{ZP3}$-expressing cells co-expressed $\text{DDX4}$ (Fig. 3). Pairwise gene analysis identified 12 $\text{DDX4}$/$\text{FIGLA}$-expressing cells, 8 $\text{DDX4}$/$\text{OOSP2}$-expressing cells, 6 $\text{DDX4}$/$\text{GDF9}$-expressing cells, and 10 $\text{DDX4}$/$\text{ZP3}$-expressing cells (Fig. 3). Thus, out of the 4 marker genes used by Wagner et al. to identify oocytes, the only gene co-expressed in all $\text{DDX4}$-expressing cells in this cluster was $\text{FIGLA}$, which is expressed in both pre- and post-miotic germ cells. A parallel analysis of $\text{NOBOX}$, which in mouse and human ovaries is robustly expressed in oocytes throughout development from the primordial follicle to metaphase-II egg stage, failed to identify a single $\text{NOBOX}$-expressing cell in the germ cell/oocyte cluster (Fig. 3). Likewise, there were no $\text{ZP1}$-expressing cells, and only 2 $\text{ZP2}$-expressing cells, identified in this cluster of cells containing candidate oocytes (Fig. 3). Of the 2 $\text{ZP2}$-expressing cells, only 1 co-expressed the 4-gene marker panel used by Wagner et al. to identify oocytes, whereas the other co-expressed $\text{FIGLA}$, $\text{OOSP2}$, and $\text{ZP3}$, but not $\text{GDF9}$ (data not shown).

Analysis of Human Unsorted Ovarian Cortical Cells for Evidence of Non–Oocyte Germ Cells

We next moved to the analysis of genes known to be expressed in primitive germ cells (see Supplementary Table S5 for more details). We identified 1 cell in the germ cell/oocyte cluster with the expression of all 5 germline genes analyzed ($\text{PRDM1}$, $\text{DPPA3}$, $\text{IFITM3}$, $\text{TUBB8}$, and $\text{DDX4}$), and 2 additional cells with expression of $\text{DPPA3}$, $\text{IFITM3}$, $\text{TUBB8}$, and $\text{DDX4}$ but lacking detectable $\text{PRDM1}$ (Fig. 4). All 3 cells localized to the same cluster of 62 cells which contained the 10 $\text{FIGLA}$/$\text{OOSP2}$/$\text{GDF9}$/$\text{ZP3}$-expressing cells. However, these 3 cells were clearly distinct from the 10 cells classified as oocytes (Fig. 4). In the 2 non–oocyte germ cells lacking $\text{PRDM1}$...
expression, we detected expression of SYCP3 (Table 1), which is required for the progression of germ cells through the early stages of the first meiotic cell division. This observation prompted us to explore additional genes involved in the early stages of meiosis-I. From this, we found that both SYCP3-expressing germ cells co-expressed STAG3 and SMC3,

**Figure 1.** Clustering and analysis of unsorted adult human ovarian cortical cells following scRNA-seq. (A) Identification of a total of 12,020 cells that formed 6 clusters following analysis of the dataset using Cell Ranger v2. (B) Identification of a total of 27,376 cells that formed 8 clusters following reassessment of the same dataset with Cell Ranger v3. (C) Cluster dendrogram depicting the lineage relationships between the 8 clusters identified using Cell Ranger v3. (D) Scatterplot analysis of the 62-cell germ cell/oocyte cluster identified using Cell Ranger v3, showing OSC gene expression scores plotted against oocyte gene expression scores. Two cells with very high OSC gene expression scores and an oocyte gene expression score of zero are highlighted by black arrows. (E) Identification of a total of 27,710 cells that formed 9 clusters following reassessment of the dataset with Cell Ranger v6. (F) Loupe Browser analysis of the Cell Ranger v6 output data, with the germ cell/oocyte cluster highlighted by the expanded box. Of the 62 cells in this cluster (each cell is depicted as an individual dot), 10 cells were identified as positive for co-expression of FIGLA, OOSP2, GDF9, and ZP3 (purple dots; examples are highlighted by black arrows) whereas 52 cells did not show co-expression of this 4-gene oocyte marker panel (light-gray dots; examples are highlighted by open arrowheads).
1 of the SYCP3/STAG3/SMC3-expressing germ cells also co-expressed SMC1a (Table 1). Notably, the proteins encoded by STAG3, SMC1a, and SMC3 are all meiosis-specific cohesin complex components involved in the formation of axial elements and cohesion of sister chromatids during meiotic prophase-I.45-48 We also identified 2 SYCP3/STAG3/SMC3-expressing cells in this cluster with co-expression of STRA8 (Supplementary Fig. S3), the latter of which is considered a critical early gene for meiosis-I progression in germ cells of both sexes.49

It is worth noting that FIGLA was detected in all 3 DPPA3/IFITM3/TUBB8/DDX4-expressing cells (Table 1). However, since these cells were distinct from the FIGLA/OOSP2/GDF9/ZP3-expressing cells (viz. candidate oocytes), the presence of FIGLA, which is not oocyte-specific,37 is still aligned with these 3 cells being classified as non-oocyte germ cells. Expression of OOSP2 was detected in the 2 DPPA3/IFITM3/TUBB8/DDX4-expressing cells, but not in the single PRDM1/DPPA3/IFITM3/TUBB8/DDX4-expressing cell (Table 1). While OOSP gene family members were first identified as encoding oocyte-enriched proteins in the mouse ovary,50,51 lineage specificity of OOSP2 in human ovaries has not been evaluated to date, and transcriptomic expression of the gene in humans is not restricted to oocytes.52

Continuing with our analysis, GDF9 was not detected in any of the DPPA3/IFITM3/TUBB8/DDX4-expressing cells found in this cluster, whereas ZP3 was detected in all 3 cells (Table 1). However, ZP3 expression was far more ubiquitous than expected, in that a total of 567 cells with ZP3 expression were identified across the population of 27 710 cells called in this dataset (Supplementary Fig. S4). Strikingly, 525 of these ZP3-expressing cells were localized outside of the germ cell/oocyte cluster (see Supplementary material, Results 1 for additional details). This widespread detection of ZP3 expression across clusters representing different lineages, most of which are somatic, is consistent with the reported low-level expression of this gene in diverse cell types and tissues in humans.52

To further assess the promiscuous nature of ZP3 expression
outside of oocytes, we analyzed a different scRNA-seq dataset derived from adult human ovarian medullary tissue.\textsuperscript{53} We did not identify a single cell with co-expression of the 4-gene marker profile used by Wagner et al.\textsuperscript{25} to identify oocytes; however, parallel analysis of this dataset identified 673 cells expressing \textit{ZP3}, again distributed randomly across the various clusters (data not shown).

Finally, given that Wagner et al.\textsuperscript{25} used \textit{DAZL} as 1 of their 3 genes for OSC screening, we identified 20 \textit{DAZL}-expressing cells in the entire dataset of 27,710 cells called using Cell Ranger v6, 10 of which were localized to the germ cell/oocyte cluster. Of these 10 cells, 5 co-expressed the 4-gene marker panel used by Wagner et al.\textsuperscript{25} to identify oocytes (Supplementary Fig. S5), consistent with past studies establishing expression and function of \textit{DAZL} in both pre- and post-meiotic germ cells.\textsuperscript{54,55} Breaking the oocyte marker panel down further, we identified 8 \textit{DAZL}/\textit{FIGLA}-expressing cells, 8 \textit{DAZL}/\textit{OOSP2}-expressing cells, 5 \textit{DAZL}/\textit{GDF9}-expressing cells, 5 \textit{DAZL}/\textit{ZP3}-expressing cells, and 5 \textit{DAZL}/\textit{GDF9}/\textit{ZP3}-expressing cells in the germ cell/oocyte cluster (data now shown).

**Analysis of Cells Sorted from Human Ovarian Cortical Tissue Biopsies using Flow Cytometry**

In the Wagner et al.\textsuperscript{25} study, the authors also reported that DDX4 antibody-based FACS, a method used by many others to specifically sort OSCs across species since 2009\textsuperscript{7-13,20,22,23,28-30} (Supplementary Table S1), led to the isolation of PVCs and not OSCs. While initially puzzling, we noted that their flow cytometry was performed using AF594 detected with a 561-nm laser in the red channel, which is widely known to detect autofluorescence as a “positive” event during FACS. In evaluating the antibody validation and gating strategies shown in the Supplementary data of Wagner et al.\textsuperscript{25}, we...
observed that the area above the cutoff designating the negative versus positive fractions in their negative control sample lacking antibody contained positive events, which represent autofluorescence. With this information in mind, PVCs are known to express autofluorescent biomolecules, such as collagen and elastin, which produce widely known artifacts in flow cytometry.\(^{56,57}\) We therefore sorted dispersed ovarian cortical tissue with a 561-nm laser in the red channel following the parameters published by Wagner et al.\(^{25}\) Using adult bovine ovarian cortical tissue for validation, a distinct population of autofluorescent events was obtained (Fig. 5).

Three-quarters of these cells were positive for SMA or CD31, which respectively mark the 2 cell types that comprise PVCs: vascular smooth muscle cells and pericytes. A distinct population of autofluorescent events was similarly detected in dispersed ovarian cortical tissue of reproductive-age women. Likewise, almost two-thirds of these events were identified as being SMA- or CD31-positive (Fig. 5). Moreover, these autofluorescent events were detectable in dispersed cell preparations from human ovarian cortical tissue irrespective of whether the samples were gated versus FSC-A or SSC-A, or if PE-Texas Red-A was plotted against a different laser (APC-A) (Supplementary Fig. S6).

### Germ Cells and PVCs Segregate into Distinct Clusters

We then dug deeper into the conclusion of Wagner et al.\(^{25}\) that we and others have mistakenly worked with PVCs in studies that have explored the characteristic germline properties, and potential clinical utility, of human OSCs\(^7,15,22,23\) (Supplementary Tables S1 and S2). Using the optimized scRNA-seq workflow described above, we identified a cluster comprised of 3310 total cells, 479 of which had the 5-gene expression profile used by Wagner et al.\(^{25}\) to identify PVCs (\(RGS5, MCAM, MYH11, RERGL,\) and \(TALGL\) (Fig. 6A). None of the cells comprising this cluster co-expressed the gene panel which identified the 3 non-oocyte germ cells (\(DPPA3, IFITM3, TUBB8,\) without or with \(PRDM1\); data not shown), and only 3 of the 3310 total cells in this cluster expressed \(DPPA3\) (Fig. 6B). Moreover, none of the 479 cells identified as PVCs using the 5-gene profile of Wagner et al.\(^{25}\) co-expressed \(DPPA3\) (Fig. 6B). Likewise, none of the 3 non-oocyte germ cells expressed the 5-gene profile used by Wagner et al.\(^{25}\) to cluster PVCs (Table 1), which led to the expected segregation of these 2 cell types into distinct and non-overlapping clusters (Fig. 6C). In fact, of the 5 genes used by Wagner et al.\(^{25}\) to cluster PVCs, \(TALGL\) was the only gene identified through individual gene-by-gene analysis to be co-expressed in the 3 non-oocyte germ cells (\(DPPA3, IFITM3, TUBB8,\) and \(DPPA3, IFITM3, TUBB8/PRDM1\)). However, \(TALGL\) expression is not specific to any single cell type, and thus its utility as a lineage marker must be viewed in context with other genes as a profile associated with a given cell type. Supporting this statement, we found that 16 291 cells of the 27 710 total cells called expressed \(TALGL\) (Fig. 6C). This included 23 cells in the 62-cell germ cell/oocyte cluster, 4 of which co-expressed...
the 4-gene profile used by Wagner et al.\textsuperscript{25} to identify oocytes (Fig. 6C).

**Discussion**

While scRNA-seq is useful as a tool to gain insights into cell lineage heterogeneity within a sample,\textsuperscript{58} a major caveat of this approach is that its failure to detect gene expression-based evidence of a given cell type after clustering analysis does not, by default, equate to that cell type being absent in the sample analyzed. This is especially apropos in attempts to identify either very rare cells or low-expression cells in a dispersed cell preparation that is heterogenous in nature, highlighting the challenges associated with the detection of stem cells in

**Figure 5.** Flow cytometric detection, isolation, and characterization of autofluorescent events in adult cow and human ovarian cortical tissue. (A–D) Representative gating strategy for doublet discrimination (forward-scatter or FSC-A; B; side-scatter or SSC-A; C) and for dead cell exclusion using 4’,6-diamidino-2-phenylindole (DAPI) labeling (D). (E, F) Comparison of autofluorescent events detected in the APC-A far-red channel (640-nm laser; E) versus the PE-Texas red-A channel (561-nm laser; F). (G–K) Autofluorescent events detected in the PE-Texas red-A channel were collected, fixed and permeabilized (G and H), and then incubated with APC-conjugated primary antibodies against SMA (Abcam ab5694) or CD31 (Invitrogen MA3100) (I and J) for determination of the total percentage of autofluorescent events that were positive for expression of either PVC marker in bovine and human ovarian cortical tissue samples (K). Data shown in (K) are the mean $\pm$ SE; $n = 4$ (CD31) or 7 (SMA), and $n = 5$ (SMA) or 6 (CD31), for bovine and human sample analysis, respectively.
tissues by scRNA-seq. The analytical pipeline used also has a significant impact on the depth and accuracy of the data obtained, especially if the objective is to produce a comprehensive snapshot of as many cells, and as many cell types, as possible in the sample analyzed. In the Wagner et al. study, their attempts to identify OSCs in a pool of 12,160 total cells called, given the extreme rarity of OSCs in adult ovaries, would be difficult even under the best of conditions (see Supplementary material Discussion 1 for additional details). Our rigorous reassessment of their unsorted cell dataset following empirical testing of numerous variables that affect the depth, resolution, and accuracy of scRNA-seq highlight how multiple decisions made by these authors for their analysis of unsorted cells actually minimized, rather than optimized, the probability of detecting rare or low-expression cell types such as OSCs. In fact, several other ovarian stem cell types were also missed by Wagner et al., including pluripotent embryonic stem cell (ESC)-like cells, mesenchymal stem cells (MSCs), and very small embryonic-like stem cells (VSELs) (Supplementary Table S7; see also Supplementary material Discussion 2), the latter of which have been reported to support postnatal oogenesis in mammalian ovaries.

These challenges were further complicated by the fact that Wagner et al. restricted their efforts to find evidence of OSCs in their entire dataset of 12,180 cells to only 15 cells that were manually selected by these authors based on the required presence of DDX4 mRNA to establish the only “cluster” of cells that could contain OSCs. The a priori assumption that all candidate OSCs must have detectable DDX4 expression using scRNA-seq is fraught with interpretational problems. For example, we found that only 5 of the 10 cells identified as oocytes using the 4-gene profile reported by Wagner et al. co-expressed DDX4. Based on their reasoning, such an approach would have removed the remaining 5 FIGLA/OOSP2/GDF9/ZP3-expressing cells lacking detectable DDX4 mRNA from further consideration as oocytes. Likewise, we could not identify a single NOBOX-expressing cell in the germ cell/oocyte cluster, even though NOBOX is highly expressed in oocytes at all developmental stages. If one evaluated this dataset for evidence of oocytes based solely on NOBOX expression, or manually created a “cluster” of NOBOX-expressing cells as the sole cell population in which any potential oocytes would be found, the reasoning of Wagner et al. with OSCs would lead to the erroneous conclusion that oocytes do not exist in adult human ovarian cortex. At the other end of the spectrum, our evaluation of ZP3, which is widely used as an “oocyte-specific” marker, revealed low-level but widespread expression of this gene across all cell clusters, most of which are somatic in origin. Thus, scRNA-seq workflow design, and any
conclusions drawn, based on the expression of a single gene being detected or not in a cell of interest lack scientific rigor and confidence.

However, optimization of the scRNA-seq workflow pipeline using a version of Cell Ranger (v3), which was available to, and used by, Wagner et al. at the time of their study but was not used for their unsorted cell analysis, allowed us to identify rare cells in their adult human unsorted ovarian cortical cell dataset with a gene expression profile that closely aligns with that of primitive germ cells, such as embryonic PGCs and adult ovary-derived OSCs. Further analysis showed that 2 of these non-oocyte germ cells expressed multiple markers of meiosis-I commitment and progression. These observations, which offer evidence of ongoing de-novo oogenesis in adult human ovaries under normal physiological conditions, are consistent with prior studies with mice which have demonstrated that resident germ cells routinely undergo meiosis in adult ovaries. Interpretational caution must still be exercised here, however, since gene expression profiling does not offer definitive evidence of the existence, or not, of human OSCs or of active meiotic entry. In this regard, all scientific studies of isolated human OSCs published to date have characterized the cells, following isolation, by gene expression profiling along with various downstream functional tests of meiotic cell division capability and/or oocyte-forming potential. Like other gene expression-based studies, the inability of scRNA-seq as a standalone approach to offer any type of functional verification of suspected lineage identity is another caveat of the use of this type of “big-data” technology and the interpretations drawn from it.

Moving on to the FACS analysis of cells identified by Wagner et al. as DDX4-positive (+) or DDX4-negative (−), the sorting strategy for OSC isolation relies on the detection of an externalized region of the C terminus of DDX4 exposed on the outside of viable cells and not simply DDX4 expression. Dual-antigen single-protein sorting studies conducted almost 10 years ago showed that OSCs could be sorted as viable cells using C-terminal, but not N-terminal, DDX4 antibodies, noting that both antibodies recognize DDX4 in oocytes in fixed ovarian sections. In turn, the viable cells sorted with the C-terminal antibody show a near-complete population shift by FACS when the same cells are permeabilized and then analyzed with an N-terminal DDX4 antibody, verifying the specificity of the sorting protocol for detection of externalized DDX4. It has also been shown that proper tissue dispersion is a crucial step to achieve viable cell isolation and to release OSCs as single cells. Human ovary tissue is particularly fibrous and difficult to disaggregate, and thus extreme care must be exercised during disaggregation to maintain cell viability. If this is not done, the possibility of non-specific antibody binding is markedly increased, which may explain why the yield of “DDX4+” events obtained by Wagner et al. with the Abcam DDX4 antibody was 3.0-6.5-fold higher than the yield of human OSCs reportedly previously using the same sorting approach with the same antibody.

Putting potential technical issues aside, it is important to emphasize that no other study that has used DDX4 antibody-based sorting to isolate OSCs, an approach first reported over 10 years ago with more than 30 corroborating studies since then, has retrieved PVCs. In addition, DDX4 antibodies have been used to sort PGC-like cells from cultures of human embryonic stem cells and induced pluripotent stem cells indicating that the utility of this approach to specifically isolate primitive germ cells is not unique to OSCs. Notably, only PGC-like cells were obtained after DDX4 antibody–based sorting in these 2 studies, even though PVCs also arise in cultures of differentiating human pluripotent stem cells. This discordance in what Wagner et al. reported regarding their isolation of PVCs instead of OSCs by this approach also extends to their own previously published findings, in which identical claims were made that human OSCs do not exist and that the sorting strategy for OSC isolation using DDX4 antibodies does not work. While those claims were experimentally disputed in this earlier study the authors similarly performed scRNA-seq on “DDX4+” cells obtained from human ovarian cortical biopsies. Their analysis in that prior study did not, however, identify PVCs as the primary cell type retrieved by FACS. Instead, out of a randomly selected population of 41 “DDX4+” cells, their gene expression associations identified a mixed population of very diverse cell types. The inconsistent outcomes reported by these authors when using DDX4 antibodies for cell sorting in their own studies may help explain why their findings diverge widely from what many others have consistently reported using the same cell sorting strategy since 2009 (Supplementary Table S1) (see also Supplementary material Discussion 1).

With this information as a preface, we designed experiments to determine how PVCs could be erroneously isolated as cells perceived to be antibody-positive using flow cytometry. Our data presented herein offer at least a reasonable explanation for this outcome, which accounts for the inherent autofluorescence of PVCs being detected as a false-positive signal during FACS. This would lead to an artifactual enrichment of these cells rather than true antibody-positive cells. In turn, our analysis of PVCs and non-oocyte germ cells in the Wagner et al. dataset demonstrated that these 2 cell types, not surprisingly, cluster separately and exhibit no overlap in gene expression profiles associated with each cell type. Thus, even if PVCs were isolated by DDX4 antibody–based FACS for reasons unrelated to endogenous autofluorescence, any downstream analysis of these cells would generate data that differ considerably from the published results from many other groups that have successfully sorted human OSCs for characterization of their germline identity and oocyte-forming properties.

Conclusions

Since the initial report on OSCs almost 2 decades ago, over 80 primary research studies have been published supporting the existence of OSCs and/or postnatal oogenesis in mammals (Supplementary Tables S1–S3). More than 30 of these have sorted OSCs from ovaries with polyclonal or monoclonal antibodies directed against the C terminus of DDX4 for in-depth characterization (Supplementary Table S1). In this same time frame, only 10 primary research papers have been published disputing the existence of OSCs and/or the occurrence of postnatal oogenesis in mammals (Supplementary Tables S1 and S3), and only 4 of these studies claimed that DDX4 antibody-based sorting fails to isolate OSCs (Supplementary Table S1; see also Supplementary material Discussion 3). With respect to human OSCs, at least 6 different groups have established, and independently corroborated, that extracellular DDX4-positive cells sorted from adult
human ovarian cortex express primitive germ lineage (but not oocyte) markers, can be expanded in number in culture, activate meiosis, and generate oocyte-like cells in vitro and oocytes in ovarian tissue.\textsuperscript{7-15} Aside from the fact that these outcomes are fully consistent with a large body of work on OSCs in other species,\textsuperscript{6,18,70} none of these endpoints are features of PVCs.

In consideration of this, along with the experimental evidence presented herein, a more reasonable conclusion from the Wagner et al.\textsuperscript{25} study is that the scRNA-seq workflow used by the authors was not designed appropriately to identify candidate OSCs, or in fact any stem cell type, in their sample. When the analytical workflow was optimized and applied to all cells of their sample equally, we uncovered evidence in their dataset of the existence of both OSCs and primitive germ cells entering meiosis I. Likewise, a more reasonable conclusion from their flow cytometry work is that these authors have had recurrent technical difficulties with FACS over the years\textsuperscript{26,68} in achieving what more than 30 other studies have already reported with respect to the sorting of OSCs from adult ovarian tissue for in vitro and in vivo characterization (Supplementary Table S1). This alternative conclusion would also remove the erroneous inference made by Wagner et al.\textsuperscript{25} that numerous other groups have mistakenly worked with PVCs, and not germ cells, in the many reports of OSCs\textsuperscript{6} (Supplementary Table S1) or PGCL-like cells\textsuperscript{64,65} published to date using DDX4 antibody-based sorting to isolate primitive germ lineage cells.

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Conflict of Interest

H.A., T.B., Y.T., O.I., H.S., R.A.A., and E.E.T.: declare no potential conflict of interests. Z.F.: declares leadership position with Foundation Medicine, Abbvie and ownership interest with Roche, Abbvie. D.C.W.: declares interest in intellectual property described in U.S. Patent 8,642,329, U.S. Patent 8,647,869, U.S. Patent 9,150,830, and U.S. Patent 10,525,086. J.L.T. declares interest in intellectual property described in U.S. Patent 7,195,775, U.S. Patent 7,850,984, U.S. Patent 7,955,846, U.S. Patent 8,642,329, U.S. Patent 8,647,869, U.S. Patent 8,652,840, U.S. Patent 9,150,830, U.S. Patent 9,267,111, U.S. Patent 9,845,482, and U.S. Patent 10,525,086.

Author Contributions

H.A., Z.F., and D.C.W: performed the analysis of the scRNA-seq dataset; H.A.: performed the flow cytometric analysis; T.B.: assisted with the experiments and performed analysis of published studies of OSCs, postnatal oogenesis, and other ovarian stem cell types; R.A.A., E.E.T. Y.T., O.I., and H.S.: collected and cryopreserved human ovarian cortical tissue for analysis; D.C.W. and J.L.T.: directed the experiments; and J.L.T.: wrote the manuscript; all authors approved the results and the final manuscript for submission.

Data Availability

The data underlying this article are available in the article and in its supplementary material.

Supplementary Material

Supplementary material is available at Stem Cells online.

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