GFP-Tagged Protein Detection by Electron Microscopy Using a GBP-APEX Tool in Drosophila

Fred Bernard1*, Julie Jouette1†, Catherine Durieu2, Rémi Le Borgne2, Antoine Guichet1* and Sandra Claret1*

1 Polarity and Morphogenesis Team, Institut Jacques Monod, CNRS, UMR 7592, University of Paris, Paris, France,
2 Imagoseine Platform, Institut Jacques Monod, CNRS, UMR 7592, University of Paris, Paris, France

In cell biology, detection of protein subcellular localizations is often achieved by optical microscopy techniques and more rarely by electron microscopy (EM) despite the greater resolution offered by EM. One of the possible reasons was that protein detection by EM required specific antibodies whereas this need could be circumvented by using fluorescently-tagged proteins in optical microscopy approaches. Recently, the description of a genetically encodable EM tag, the engineered ascorbate peroxidase (APEX), whose activity can be monitored by electron-dense DAB precipitates, has widened the possibilities of specific protein detection in EM. However, this technique still requires the generation of new molecular constructions. Thus, we decided to develop a versatile method that would take advantage of the numerous GFP-tagged proteins already existing and create a tool combining a nanobody anti-GFP (GBP) with APEX. This GBP-APEX tool allows a simple and efficient detection of any GFP fusion proteins without the needs of specific antibodies nor the generation of additional constructions. We have shown the feasibility and efficiency of this method to detect various proteins in Drosophila ovarian follicles such as nuclear proteins, proteins associated with endocytic vesicles, plasma membranes or nuclear envelopes. Lastly, we expressed this tool in Drosophila with the UAS/GAL4 system that enables spatiotemporal control of the protein detection.

Keywords: APEX, nanobody, green fluorescent protein, ovarian follicle, electronic microscopy, GBP, Drosophila melanogaster

INTRODUCTION

In cell biology studies, protein localization is crucial to understand the cellular functions of proteins and for understanding the dysfunction of proteins in diseases. For years, the technique used for this purpose was immunohistochemistry. It requires specific antibodies directed against each of the proteins of interest (POI). However, the production of good quality primary antibodies is random and labor-intensive. Once obtained, it remains a resource with limited availability.

The advent of genetically targetable fluorescent protein tags has offered a possibility to bypass the requirement of antibody production against each POI. In addition, fluorescent tags have further expanded the field of possibilities to in vivo localization in living tissues or cells. Therefore, in
Drosophila, where large scale projects are regularly conducted, various programs and consortiums have developed systematic approaches with the objective of creating lines expressing a fluorescent version of each protein of the proteome. Different approaches have been used to generate protein trap lines where an artificial exon encoding GFP is inserted into the genome (Morin et al., 2001; Clyne et al., 2004; Kelso et al., 2004; Buszczak et al., 2007; Quiñones-Coello et al., 2007; Lowe et al., 2014; Nagarkar-Jaiswal et al., 2015). Currently, the CRISPR technique better facilitates the creation of fusion proteins regulated by their endogenous environment, thereby the number of fluorescently tagged proteins generated by individual labs continuously increases.

In the vast majority of studies, the experiments described above are performed using light microscopes, which are fast, cheap and simple. The conventional fluorescence microscopy has a spatial resolution within a 200-300 nm range and it reaches a maximum of 10 nm in super-resolution microscopy but requires specialized equipment and/or fluorophores. In these conditions, intracellular localization often requires the co-localization with a fluorescent marker of organelles or compartments, although the size of many organelles is below the resolution limit of these microscopes.

Due to the imprecision of this approach, a high-resolution analysis becomes necessary through, for example, electron microscopy (EM)-based detection. Although EM achieves much higher spatial resolution (~1 nm in biological samples), the localization of proteins by EM approaches remains rare. Several reasons lead to this situation. High quality results by EM immunolocalizations are difficult to obtain. Indeed, when performed on whole tissue, immunolocalization protocols include permeabilization steps that degrade intracellular structures. Alternatively, immunolocalizations performed on ultra-thin sections have only little epitope accessible to antibodies (Sosinsky et al., 2007; Schnell et al., 2012). Moreover, contrasting agents have a negative impact on the antigen-antibody binding, therefore protocols aim at maintaining them at low levels. This leads to images with poor contrast and makes the subsequent identification of ultrastructures difficult. Another alternative is then to perform ultrathin sections of cryoprotected samples infiltrated by 2.3 M sucrose followed by an immunolabeling of the ultrastructure of the sample. APEX has been used as a tag in many studies and has largely proven its efficiency, making it now a tag of choice for the detection of fusion proteins in EM (Martell et al., 2012, 2017; Ariotti et al., 2015; Chen et al., 2015; Lee et al., 2016; Lin et al., 2016; Ludwig, 2020; Tan et al., 2020).

Here we have chosen to use the APEX2, a version of APEX (K14D, W41F, and E112K) with an additional mutation (A134P) (Lam et al., 2014). It has the same advantages of APEX while producing the DAB polymer with faster kinetics and incorporating the heme cofactor more efficiently. Similarly to an approach that has been reported to work successfully in zebrafish (Ariotti et al., 2015), we have created a GBP-APEX2 tool that combines the APEX2 with a GFP binding protein (GBP). The GBP corresponds to the coding sequence of an anti-GFP nanobody which is a single-domain polypeptide derived from the variable heavy chain (Vhh) of the heavy chain-only antibodies of camelids (Kirchhofer et al., 2010; Kubala et al., 2010). This GBP domain will allow the association of APEX with any GFP-tagged protein in vivo. In previous studies, it has been shown that individual intermediate filaments can be resolved (Ariotti et al., 2015) indicating that APEX-GBP allows a spatial resolution of ~10 nm.

The APEX-GBP strategy avoids the need to create new APEX-tagged transgenic lines for each new POI, and takes advantage of all the GFP-tagged lines already available in the Drosophila community. In addition, to further increase the adaptability and the versatility of our tool, we placed this fusion construct under the regulatory region of UAS sequences. These sequences trigger the expression of coding sequences placed downstream, when they are bound by the GAL4 transcription factor. Numerous Drosophila strains, with different expression patterns of GAL4 are available and thus, spatiotemporal control of expression can be achieved (Brand and Perrimon, 1993; Duffy, 2002).

As a proof of principle, we present here the localization of several GFP-tagged proteins and describe a detailed protocol applicable to the Drosophila ovarian follicle (Figure 1).

### MATERIALS AND EQUIPMENT

**Preliminary Notes**

Many of the chemicals used during the APEX reaction and EM steps are hazardous to humans and the environment. Therefore, pay attention to the attached Material Safety Data Sheets and handle these reagents with care: glutaraldehyde (toxic), paraformaldehyde (carcinogen, toxic), DAB (carcinogen), cacodylate buffer (toxic, arsenic), hydrogen peroxide (corrosive), uranyl acetate (radioactive), lead citrate (toxic) and osmium tetroxide (highly volatile, toxic and highly reactive). Wearing gloves as well as adapted personal protective equipment and manipulating under a fume hood are essential.

**Fly Stocks**

Fly strains and crosses were raised on standard cornmeal food at 25°C. To overexpress UAS transgenes specifically in germline cells, nos-GAL4VP16 [P(mw, GAL4:VP16-nos.UTR) CG6325(MVD1)] (DGRC Kyoto #107955) and mat-Tub-Gal4VP16 [P(mw, mat-alpha4-Gal4-VP16)]...
FIGURE 1 | Ovariole organized in a succession of developing follicle. Nuclear envelopes (red) are stained with alexa594-WGA and the cortical actin (green) revealed with Alexa448-Phalloidin. Follicles are developmental subunits in which the oocyte (Oo) develops. The oocyte is located at the posterior of the follicle and is associated with 15 additional germ cells named nurse cells (NC). A monolayer of follicular cells (FC) surrounds the germline. On the example presented here, the ovariole exhibits at its anterior extremity (left) a region called germarium in which the stem cells proliferate. At the other extremity (right) a stage 8 showing an oocyte that represents almost one half of the follicle.

(Januschke et al., 2002) were used. To overexpress UAS transgenes in somatic follicular cells, Tj-GAL4 [y,w; P(GawB)NP1624] (DGRC Kyoto #104055) was used. The strains UASp-APX2-GBP and UASp-6myc-APX2-GBP are from this study. The following GFP strains were used: P(BazBAC.GFP) aka P[w + , FRT9-2]18E, f. baz [815.8], P[CaryP, PB[BAC Baz-sfGFP2]attP18] (Besson et al., 2015); P(mudBAC.GFP) (Bosveld et al., 2016); UASp-Baz-GFP (Benton and St Johnston, 2003); UASp-Rab5-GFP (Dong and Wu, 2013); hsp-flp; FRT79D ubi-nlsGFP (gift from JR Huynh); Tub-GFP-Rab6 (Januschke et al., 2007), RanBP2-GFP (Hampoolz et al., 2019).

Generation of Transgenic Flies
APX2-GBP from pCSDEST2 APEX2-GBP (Plasmid #67651, Addgene) was subcloned in pENTR™/D-Topo. Using the Gateway™ recombination cloning, APEX2-GBP sequence was inserted in pPMW (promotor UASp with a N-terminal 6myc tag), in pPW (promotor UASp without tag) from the Drosophila Gateway™ Collection. The transgenic flies have been generated by random insertion by the BestGene Company (United States).

Reagents and Equipment for Ovary Dissection and Immunostaining
- Bovine Serum Albumin (Thermo Fisher Scientific, BP1600).
- Chicken Anti-APX2 antibody (Innovagen PA-APX2-100) raised against AA126-146.
- Mouse anti-Myc/c-Myc 9E10 antibody (SantaCruz, sc40).
- Anti-Mouse Secondary antibody, Alexa Fluor 546.
- Goat anti-Chicken IgY (H + L) Secondary Antibody, Alexa Fluor 546.
- Phosphate buffered saline pH 7.4 (PBS).
- Tween 20 (Sigma-Aldrich, P1379-1L).
- Triton X-100 (Sigma-Aldrich, T8787-100ML).
- Paraformaldehyde 16% (w/v) in sealed 10 mL glass ampules (Avantor, 43368.9L).
- Citifluor™ Mountant Solution AF1 (Electron Microscopy Sciences, 17970-25).
- Forceps Dumont #5 (Carl Roth, K342.1).
- Stainless steel needles (Entosphinx, 20).
- Colorimetric 8 cell tray (Kartell Labware, 357).

Reagents and Equipment for EM Sample Preparation and Detection
- Methylene blue staining solution (Richardson et al., 1960) (Methylene blue 0.5%, azur II 0.5%, Sodium borate 0.5%).
- Glass microscopy slides (Fisher Scientific, 1018049).
- Glutaraldehyde EM grade 25%, in sealed 10 mL glass ampules (EMS 16220).
- 3,3′-Diaminobenzidine tetrahydrochloride (DAB; Sigma-Aldrich, D5905).
- Hydrogen peroxide (H2O2), 3% (Boster Immunoleader AR1108).
- Sodium cacodylate buffer 0.2 M pH 7.4 (EMS 11652).
**STEPWISE PROCEDURE**

**Fly Handling**

In food vials, cross 5–10 virgin females with 3–5 males of the desired genotype and hold the vial at 25°C. After fly hatching select the females of the correct genotype, transfer them with few males in fresh food vials supplemented with dry yeast for their ovaries to fatten up and leave them for one or 2 days before dissection.

**Microdissection**

1- Anesthetize the flies on a pad, with carbon dioxide.
2- Under the dissecting microscope, pick up one female with a pair of forceps and immerse it in a large drop (50–100 μL) of PBT (PBS with 0.1% Tween 20) at room temperature.
3- Hold the fly by the thorax with one pair of tweezers, and pull the dorsal abdominal cuticle around the A4–A5 segmental boundary with another pair of forceps.
4- Isolate and detach the pair of ovaries, which can fill up to 2/3 of the female abdomen, and should be readily available upon cuticle removal.
5- Tease apart the ovarioles of each ovary. While holding the posterior end of the ovaries (older stages) with a forceps, pass a needle in between the ovarioles toward the germarium at the anterior end of the ovary.
6- Transfer the ovaries into a 2 mL centrifuge tube containing 200 μL of PBT at room temperature and continue the experiment quickly.

**Fixation and DAB Reaction**

7- Remove the PBT and add 500 μL of the fixative solution (2.5% paraformaldehyde, 1% glutaraldehyde in 0.1 M sodium cacodylate buffer).
8- Keep it 20 min at RT then move to 4°C.
9- Keep it at 4°C for 1 h, in the dark. *During this time, prepare the DAB solution.*
10- Wash three times 5 min in 0.1 M sodium cacodylate buffer at 4°C.
11- Prepare the DAB solution (1 mg/ml DAB, 0.1 M sodium cacodylate buffer). 1.5 mL per sample is required:
   - Dissolve one tablet of 10 mg DAB in 5 ml of H₂O with 5 min vigorous vortexing.

- Dilute 1:1 the DAB/H₂O with 0.2 M sodium cacodylate buffer.
- Remove undissolved precipitates with syringe filtration using a 0.2 μm filter (Millipore).
12- Add 500 μL of the final solution to the sample.
13- Allow to react for 30 min (*Increased time reduces background*).
14- Replace the solution with a DAB/Cacodylate + 5.88 mM H₂O₂ solution.
15- Incubate for 20 min at RT.
16- Stop the reaction with 3 min × 2 min washes with 0.1 M sodium cacodylate buffer.

**Post-Fixation**

17- Prepare post-fixative solution.
   - (a) 1% osmium tetroxide (prepared from 4% stock solution).
   - (b) 1.5% Potassium hexacyanoferrate(II) trihydrate (prepared from stock powder).
   - (c) 0.1 M cacodylate buffer (prepared from 0.2 M stock solution).
18- Incubate for 1 h at 4°C.
19- Wash three times 2 min in 0.1 M cacodylate buffer.
20- Wash three times 2 min in H₂O.

**Dehydratation**

21- Incubate 10 min in 30% EtOH solution.
22- Incubate 10 min in 50% EtOH solution.
23- Incubate 10 min in 70% EtOH solution.
24- Incubate 10 min in 90% EtOH solution.
25- Incubate twice 10 min in 100% EtOH solution.

**Resin**

26- Incubate in resin LV agar/EtOH (1/1) overnight.
27- Incubate twice in resin for 1 h.
28- Mount the samples between two sheets of fluoropolymer film, separated by a fluoropolymer film spacer. This step is important because it allows the ovarioles to be laid out flat in order to select the right stage of development and to orientate them.

28.1: Three pieces of fluoropolymer film embedding film are cut in the dimensions of a microscopy slide (75 mm × 25 mm).
28.2: In the center of one of the three pieces of fluoropolymer film a square of 20 mm × 20 mm is cut out.
28.3: Place the first piece of fluoropolymer film on a microscopy slide.
28.4: Superimpose the hollowed film.
28.5: Pipet the samples in resin.
28.6: Spread out the ovarioles.
28.7: Carefully apply the third piece of fluoropolymer film to minimizing the formation of bubbles.
28.8: Put a microscopy slide on top to make a sandwich.
be treated according to the following protocol: the localization of the GFP-tagged protein. This can be verified in the tissue Co-localization by Fluorescent Microscopy

Validating APEX2-GBP and GFP Expressions and the sample preparation.

Validating APEX2 Activity by Light Microscopy

A critical step in this protocol is the ability of APEX2 to convert the DAB into a polymer. The polymer produced by the APEX enzymes is osmiophilic and thus can be visualized in EM, but it can also be visualized using light microscopy appearing as light brown stain (Figures 3A,B). Usually, the subcellular localization of APEX in the tissue can be roughly distinguished and bodes well for visualization in transmission EM (TEM). As an illustration, we decided to detect Baz-GFP with the GBP-APEX2 construct and look at the DAB product with a transmission light microscope. When the GBP-APEX2 is specifically expressed in the germline, under the regulatory sequences of nanos-GAL4 (nos-GAL4VP16), a brown precipitate is accumulated only in the nurse cells and the oocyte (Figure 3A). Moreover, we could clearly see a stronger accumulation at the anterior of the oocyte where PAR3/Baz is normally enriched. Alternatively, when the GBP-APEX2 is specifically driven in the follicular cells with the traffic-jam-GAL4 (tj-GAL4), we observed a strong staining in the follicular epithelium that surrounds the ovarian follicle. The brown labeling is, as expected, more intense at the apical side of the cells (Figure 3B). The light brown staining observed over the germ cells corresponds to signals accumulated in the follicular cells above them. These experiments show that the fixation procedure does not alter the enzymatic activity of the APEX, nor the specificity of the DAB precipitate accumulation in cells expressing the GBP-APEX construct.

Validating Sample Preparation

The quality of TEM preparation is a function of the correct completion of several crucial steps i.e., fixation, dehydration and embedding in a resin. The quality of embedding is very important and can be checked by the analysis of semi-thin sections of the sample under a light microscope. For thick samples like
FIGURE 2 | Detection APEX2-GFP protein by immunofluorescence. (A,B) In mud-GFP; mat-Tub-Gal4VP16, UAS-mycAPEX2-GBP ovarian follicle, the GFP signal is detected around the nucleus of the oocyte (A). With anti-myc antibodies (A’), we observe a similar pattern that overlaps with the GFP fluorescence (A”). Similarly, in nos-Gal4VP16, UAS-APEX2-GBP, UAS-Par3-GFP egg chamber, we observe the GFP (B) associated with the membranes. Anti-APEX antibodies (B’) reveal an overlapping staining (B”).

the Drosophila oocyte, this step can also be used to screen for adapted z-position before collecting ultra-thin sections for TEM. Semi-thin sections (0.4 µm thick) can be colored with methylene blue in order to better visualize cell morphology. Methylene blue staining allows one to recognize cell nuclei (Figure 3C) but also to reveal potential issues with the fixation step that could affect tissue morphology (Figure 3D).

After inclusion in resin (see above), we proceed to the following steps:

1- Semi-thin sections are generated with an ultramicrotome (Leica Microsystems UC6).
2- Two or three sections are deposited within a drop of water onto a glass slide.
3- Slides are placed 15–30 s on a preheated hot plate to dry the sections.
4- When the water is totally evaporated, apply a drop of the methylene blue staining solution onto the sections to cover the entire surface.
5- Incubate for 30 s on the hot plate.
6- Remove the staining solution by rinsing with distilled water.
7- Last traces of water are removed by placing the slide for 30 s back on the hot plate.
8- Observation under a dissecting microscope.

It should be noted that various polychromatic staining techniques can be used for embedded tissue sections (Toluidine blue, Basic fuchsin, and Malachite green).

Proof of Principles

In order to validate our APEX tool in Drosophila, we have chosen to test, in the ovarian follicle, different GFP-tagged proteins associated with various subcellular compartments. These proteins are either overexpressed with the UAS/GAL4 system or expressed under the control of their own promoters with transgenes, or directly tagged in the genome by CRISPR mediated GFP insertion. For each condition, GFP was visualized by confocal microscopy, in parallel to the treatment for TEM observation. The APEX2-GBP construction is expressed under the control of UAS promoter and nos-GAL4VP16 driver in the germline cells or tj-GAL4 driver in the follicle cells.

In a first attempt, we detected DE-Cadherin (DE-Cad) encoded by the shotgun gene in Drosophila. DE-Cad is a component of the adherens junction that localizes at the plasma membrane. Using a DE-Cad-GFP knockin line (Huang et al., 2009), we observed by fluorescence a signal at the oocyte plasma membrane with patches of higher intensity (Figures 4A,A’). Similarly, APEX detection and visualization by TEM revealed electron dense patches associated with plasma membranes (Figure 4A”).

We then addressed whether this approach is suitable to detect protein involved in cellular trafficking, such as the GTPhases RAB5 and RAB6. Fluorescence detection of the RAB5-GFP tagged protein, expressed under the control of UAS sequence, displays a cortical signal along the plasma membrane of the oocyte (Figures 4B,B’) as expected with the previously described RAB5 association with early endosomes (Zerial and McBride, 2001; Compagnon et al., 2009). With our detection method by TEM, we have observed signals associated with vesicles near the plasma membrane (Figure 4B”). Interestingly the DAB precipitate seems to be organized in nanodomains on the endosomes as it has been proposed previously (Franke et al., 2019). Concerning RAB6, this GTPase is known to be associated with medial Golgi and Trans Golgi Network (Antony et al., 1992) and it has been shown to regulate
transport between early and late Golgi compartments and to sustain Golgi morphology (White et al., 1999; Januschke et al., 2007). With fluorescence we observed a diffuse staining pattern, with a few more intense dots, scattered within the cytoplasm (Figures 4C,C’). APEX-GBP revealed by TEM highly contrasted dots in the close vicinity of Golgi apparatus, a location consistent with the previously described role of RAB6 (Figure 4C”).

To monitor the versatility of the UAS-GBP-APEX2 tool, we tested the use of this tool in the somatic cells surrounding the ovarian follicle. For this purpose, we monitored the localization of the PAR3/Baz polarity protein tagged with GFP and expressed at endogenous levels. Upon APEX detection and visualization by TEM, dense patches were easily identified at the level of the adherens junctions as expected for PAR3/Baz (Figures 5A,A’ and Supplementary Figure 1). The APEX-GBP system is therefore effective in tracking the EM localization of GFP-tagged proteins independently of their expression level.

To test, if the method is sensitive enough to reflect differences in protein quantity, we chose to follow the nuclear envelope repartition of the Mud/NuMA protein, that is known to be asymmetrically distributed at the oocyte nuclear envelope (Yu et al., 2006; Tissot et al., 2017; Figures 5B,B’). Accordingly, the ultra-thin sections observed in TEM exhibit more dense signals on the portion of the nuclear envelope facing the posterior membrane of the oocyte (Figure 5B”). DAB precipitates on the opposite side of the nucleus are much less present showing that our conditions can detect different quantities of proteins.

One limit of the APEX approach is the diffusion of the DAB precipitate formed by the enzymatic activity. We thus decided to estimate this diffusion by looking at a protein with a precise location, i.e., the RanBP2/Nup358 protein that is an outer component of nuclear pore complexes (Bernad et al., 2004). Strikingly, the diffusion observed with the detection of RanBP2-GFP is restricted outside of the nucleus according to the known location of the protein (Figure 5C). Furthermore, we do not

FIGURE 3 | Method validation. (A) Images of nos-Gal4; UAS-APEX2-GBP, UAS-Par3-GFP ovarian follicle acquired on transmission light microscope reveal DAB precipitates specifically in the germline. Stronger accumulation at the anterior of the oocyte (arrows) is coherent with Par3 distribution profile. (B) In P(BazBAC.GFP); UAS-APEX2-GBP/Tj-GAL4, DAB staining is only observed in follicular cells (arrows). Note that the brown shade over the germline staining is due to surrounding follicular cells. (C,D) Methylene blue staining reveals morphology of the follicle and allows visualization of nuclei. Thereby, we can also verify the developmental stage of the egg chamber (C). In addition, issue with fixation is also revealed by burst ovarian follicles [arrows, (D)].
FIGURE 4 | Expression profiles of DE-Cad-GFP (A–A”), Rab5-GFP (B–B”) and Rab6-GFP (C–C”), revealed by fluorescent microscopy (A,A’,B,B’,C,C’) or by electron microscopy (A”,B”,C”). DE-Cad localizes at the plasma membrane [(A), higher magnification (A’)]. DAB precipitates are visualized (arrowheads) near plasma membranes (PM) separating the oocyte and a nurse cell (A”). Rab5 that has a cortical localization revealed by fluorescence [(B), higher magnification (B’)], is detected at a small distance of the plasma membrane at the surfaces of vesicles [(B”), arrowheads)]. Rab6-GFP displays a diffuse cytoplasmic signal (C) with some dots revealed at higher magnification (C’). By electron microscopy, DAB precipitates are detected in the vicinity of Golgi apparatus [(C”), arrowhead], (m) mitochondria, brackets highlight a Golgi unit, (VM) vitelline membrane, (FC) follicular cells. Number of observations DE-Cad-GFP (n = 4), Rab5-GFP (n = 2) and Rab6-GFP (n = 6).

detect staining when large portions of NE devoid of visible nuclear pores are observed. In this case, no diffusion could be detected inside the nucleus, showing that this method is suitable to decipher if a given protein is associated with the inner or outer membrane of the NE.

This observation prompted us to test if the UAS-GBP-APEX2 tool could be used to detect protein within the nucleus, despite our choice not to include a nuclear localization sequence in our GBP-APEX construct. In order to address if a nucleus-resident protein could nevertheless be monitored in TEM using our APEX-GBP tool, we expressed a nls-GFP transgene concomitantly with APEX-GBP in the ovarian follicle (Figure 6).

In Figure 6A, the APEX-GBP is expressed only in the germline and we can observe a more intense staining in the germline nuclei (Figure 6, white N). Note that a weak staining in the germline cytoplasm can also be observed indicating the presence of APEX-GBP not associated with GFP. Importantly, in the follicle cells that serve as a control condition, the nucleus is lighter than the surrounding cytoplasm (Figures 6A,B, black n), unlike the germline where the nucleus is more strongly stained (Figures 6A,B, white N) than the cytoplasm.

DISCUSSION: CRITICAL PARAMETERS AND TROUBLESHOOTING

Driver/APEX-GBP Couple

Having a bi-partite system where the APEX is uncoupled from the POI has many advantages as mentioned previously. However, in this system APEX is expressed throughout the whole cell independently of the POI’s subcellular location and thereby induces a background signal. Therefore, it is important to perform control experiments when studying a protein for the first time (see below). This is also exemplified by the detection of signals unspecific to our protein when immunofluorescence experiments are performed with anti-APEX antibodies (see Figure 2). However, in TEM, this does not prevent an accurate detection of Mud only at the nuclear envelope of the oocyte. It
is noteworthy that negative controls display much lower global signal, indicating that a significant but acceptable level of noise is induced by this condition. We can speculate that the local concentration of APEX protein is higher when bound to the POI and thereby create a signal/noise ratio in favor of the detection.

When we used the Tj-GAL4 driver to detect the nls-GFP in the follicular cells, we observed a cytoplasmic background of a similar level to the nuclear signal associated with the nls-GFP transgene (Figure 6E). It is then difficult without control to distinguish between specific and non-specific labeling. Nonetheless, in the same cells and with the same driver, the signal of PAR3/Baz-GFP, expressed at endogenous levels with a BAC transgene, was strong enough to be unambiguously identified (Figure 5A’). These examples reveal that the driver/APEX-GBP couple has to be carefully chosen in order to maximize the signal to noise ratio. It is noteworthy that several methods have recently been reported to improve the signal-noise ratio, i.e., the use of conditionally stable nanobodies for GFP fused to APEX, that favor degradation of unbound nanobodies by the proteasome reduces background APEX signals (Ariotti et al., 2018), or the possibility to convert the oxidized diaminobenzidine reaction product of APEX into a silver/gold particle that in addition provides a readily quantifiable particulate signal (Rae et al., 2021).

Both approaches have been proven successful in cell cultures and remain to be tested and adapted in vivo to thicker tissue like Drosophila follicles.

**Negative Controls**

Transmission EM images are displayed in gray levels that reflect the density of the structure encountered by the electron beam. In order to help visualize cellular structures and increase the contrast, the samples must be incubated with osmium tetroxide, uranyl acetate and/or potassium ferricyanide. DAB precipitates also appear as dense structures, thereby it could be challenging to identify the osmiophilic precipitate produced by APEX especially when the POI has an unknown location. Therefore,
FIGURE 6 | Nuclear detection of APEX2. (A–C) In hsp-flp; FRT79D ubi-nlsGFP/nos-GAL4[16], UAS-APEX2-GBP follicles, strong DAB staining is observed in germline nuclei (N) [(A)], higher magnifications in panels (B,C)]. (D) Control sample without the APEX2-GBP transgene exhibits a strong difference between the contrasts of the nucleus that is lighter than the cytoplasm. Comparison of the contrasts between panels (C,D) clearly shows that the stronger signal in the nucleus of hsp-flp; FRT79D ubi-nlsGFP/nos-GAL4[16], UAS-APEX2-GBP ovarian follicles is specific. (E) In hsp-flp; Tj-GAL4; FRT79D ubi-nlsGFP/UAS-APEX2-GBP, nuclei of the follicular cells display strong accumulation of DAB precipitates. (N) germline nuclei, (n) follicle cells nuclei, (Y) yolk vesicle, (m) mitochondria, (al) annulate lamellae. Number of observations (A–C): (n = 5), (D) (n = 2), (E) (n = 2).
we suggest performing negative controls with samples devoid of APEX-GBP proteins (no GAL4 transgene or no UAS-APEX2-GBP transgenes) and samples lacking GFP proteins. If a staining is reproducibly observed in APEX expressing tissue and never observed in controls, we can be confident about the specificity of the staining. In the case of nls-GFP localization with APEX, we thus compared the contrasts existing in the oocyte nucleus in the presence (Figure 6C) or absence (Figure 6D) of the APEX2 transgene. We also observed APEX-related contrast in the absence of GFP-labeled proteins (Supplementary Figure 2).

**Golgi Apparatus**

In the Drosophila ovarian follicle, we often visualized an electron dense staining, independently of the presence of APEX, within the Golgi cisternae (Supplementary Figure 2). We believe that it depends on a glutathione peroxidase (PHGPx) located in the Golgi apparatus (Missirlis et al., 2003). The visualization of this staining is not fully penetrant but the APEX-GBP tool is probably not appropriate for the detection of POI located in this organelle.

**Weak APEX Signal**

$\text{H}_2\text{O}_2$ is necessary for the oxidation reaction to occur, however, it has also been reported that long incubation could inhibit the reaction (Ludwig, 2020). Therefore, in case of weak signal, the DAB labeling can not very easily be adjusted by changing its duration. Instead it has been suggested that lowering $\text{H}_2\text{O}_2$ concentration to 0.5 mM greatly enhances APEX2 activity and sensitivity, and results in an increased contrast in TEM (Ludwig, 2020). It is therefore possible that small amounts of APEX protein can be detected by adjusting the $\text{H}_2\text{O}_2$ concentration within the 10–0.5 mM range. Here we used in routine an intermediate concentration of 5.9 mM.

**Time Considerations**

Dissection, fixation and washing procedures take around 4 h and are followed by 1 h post-fixation incubation, 1 h dehydratation before an overnight incubation in the resin. Embedding takes a further 24 h, followed by an additional day required for resin polymerization. The embedded sample can be stored indefinitely before sectioning.

**CONCLUSION**

The use of APEX has recently gained momentum in the scientific community as it offers an easy, cheap and rapid way of localizing a POI with the resolution of EM. This peroxidase has already been used in Drosophila in its original version (Chen et al., 2015; Lin et al., 2016).

By coupling APEX2 to the GBP nanobody, we have created a new tool that can be used in any cell type and for any GFP (and derivatives) labeled protein in Drosophila. We show here the flexibility of this tool to identify the nanometric localization of proteins in different compartments by TEM. Notably, proteins either expressed by their endogenous promoters or over-expressed have been detected with the exact same conditions, showing that this protocol does not need too much adaptation from one POI to another. We have tested our protocol with classical TEM, but there have been reports showing that EM volume imaging such as SBF (serial block face) technology could also be successfully combined with APEX approaches (data not shown and Ariotti et al., 2015; Ludwig, 2020).

The bi-partite detection of APEX also offers the possibility for a reliable technique of correlative light and EM (CLEM) whereby the GFP-tagged POI can be visualized using fluorescence microscopy, and the DAB precipitate generated by APEX can be identified by EM at the place of the GFP-tag (For review, see Ariotti et al., 2015; Ludwig, 2020).

Finally, another popular use of APEX are the proteomic approaches. Indeed, in addition to DAB, APEX can use biotin-phenol as substrate. In presence of $\text{H}_2\text{O}_2$, APEX then catalyzes the formation of biotin-phenoxyl, which can covalently bind electron-rich amino acids such as tyrosine in the proteins located in close proximity. It is estimated that modified proteins are within a radius of 20 nm. Biotinylated proteins are subsequently identified by mass spectrometry. Several studies have successfully developed this approach including in Drosophila (Markmiller et al., 2018; Marmor-Kollet et al., 2020) and the Drosophila oocyte (Mannix et al., 2019; Gerdes et al., 2020). In addition, APEX can also biotinylate guanosine in RNA, a recent study has used this property to determine subcellular transcriptome after RNA sequencing (Fazal et al., 2019). All these studies have been performed by using direct fusion of APEX to a POI. Theoretically, our APEX-GBP tool could also be suitable for these approaches and would prevent labs from generating new constructs given all the already existing GFP-tagged proteins.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

**AUTHOR CONTRIBUTIONS**

JJ and SC designed the project and performed the clonage experiments. CD and RB performed the TEM experiments. SC and FB conducted data interpretation and writing. All authors contributed to manuscript revision, read, and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcell.2021.719582/full#supplementary-material

Supplementary Figure 1 | Immunostaining anti-APEX2 on PmudGFP; UAS–APEX2–GBP ovarian follicles. Specific but diffuse staining is revealed in follicular cells.

Supplementary Figure 2 | (A) Examples of non-specific staining independent of APEX in Golgi apparatus. (B) Negative control of APEX experiment. The APEX procedure was realized on mudGFp; UAS–APEX2–GBP in absence of APEX2–GBP, we revealed only the cell structures. (G) Golgi apparatus, (N) nuclei, (m) mitochondria, (PM) plasma membrane, (ER) endoplasmic reticulum.
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