**Drosophila Ana2 is a conserved centriole duplication factor**

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In Caenorhabditis elegans, five proteins are required for centriole duplication: SPD-2, ZYG-1, SAS-5, SAS-6, and SAS-4. Functional orthologues of all but SAS-5 have been found in other species. In Drosophila melanogaster and humans, Sak/Plk4, DSas-6/hSas-6, and DSas-4/CPAP—orthologues of ZYG-1, SAS-6, and SAS-4, respectively—are required for centriole duplication. Strikingly, all three fly proteins can induce the de novo formation of centriole-like structures when overexpressed in unfertilized eggs. Here, we find that of eight candidate duplication factors identified in cultured fly cells, only two, Ana2 and Asterless (Asl), share this ability. Asl is now known to be essential for centriole duplication in flies, but no equivalent protein has been found in worms. We show that Ana2 is the likely functional orthologue of SAS-5 and that it is also related to the vertebrate STIL/SIL protein family that has been linked to microcephaly in humans. We propose that members of the SAS-5/Ana2/STIL family of proteins are key conserved components of the centriole duplication machinery.

**Introduction**

The centriole is composed of a radial array of nine microtubule (MT) triplets, doublets, or singlets depending on species and cell type. Centrioles are required to make two important cellular structures: centrosomes and cilia. The centrosome consists of a pair of centrioles surrounded by pericentriolar material (PCM) and is the major MT organizing center in many animal cells (Rieder et al., 2001; Doxsey et al., 2005). Cilia are formed when the centriole pair migrates to the cell cortex, and the older, mother, centriole forms a basal body that nucleates the ciliary axoneme. Many different cell types possess cilia, and they have multiple roles in development (Gerdes et al., 2009).

To ensure their inheritance by each daughter cell, centrioles duplicate precisely once per cell cycle. This process must be tightly regulated. Failure in centriole duplication leads to catastrophic errors during embryogenesis in both worms and flies (O’Connell et al., 2001; Stevens et al., 2007), and an increasing number of human diseases have been linked to defects in centrosome and/or cilia function (Badano et al., 2005; Sharma et al., 2008). Centriole overduplication can be equally damaging, as excess centrioles are frequently observed in human tumors (Nigg, 2002), and there appears to be a direct causative relationship between centriole overduplication and tumorigenesis in flies (Basto et al., 2008).

In canonical centriole duplication, a new daughter centriole grows at a right angle to the mother centriole. A series of genome-wide RNAi and genetic screens in worms have found just five proteins essential for centriole duplication: SPD-2, ZYG-1, SAS-5, SAS-6, and SAS-4 (O’Connell et al., 2001; Kirkham et al., 2003; Leidel and Gönczy, 2003; Dammermann et al., 2004; Delattre et al., 2004; Kemp et al., 2004; Pelletier et al., 2004; Leidel et al., 2005). SPD-2 is required to recruit the kinase ZYG-1 to the centriole, and both proteins then recruit a complex of SAS-5 and SAS-6 (Delattre et al., 2006; Pelletier et al., 2006). SAS-5 and SAS-6 are mutually dependent for their centriolar localization and are in turn needed to recruit SAS-4 (Leidel et al., 2005).

Although DSpd-2 is not essential for centriole duplication in flies (Dix and Raff, 2007; Gianantsi et al., 2008), and no SAS-5 homologues have been identified outside worms, proteins related to ZYG-1, SAS-6, and SAS-4 have a conserved...
role in centriole duplication in other systems. In *Drosophila*, for example, the kinase Sak, which is related to ZYG-1, and the homologues of SAS-6 (DSas-6) and SAS-4 (DSas-4) are required for centriole duplication (Bettencourt-Dias et al., 2005; Basto et al., 2006; Peel et al., 2007; Rodrigues-Martins et al., 2007a). Recently, however, several additional proteins have been identified in cultured fly cells that are potentially involved in centriole duplication (Goshima et al., 2007; Dobbeltrae et al., 2008). Here, we set out to identify which of these potential duplication factors are likely to function as upstream regulators of centriole formation.

**Results and discussion**

**Ana2 and Asterless (Asl) can drive the de novo formation of centriole-like structures**

Genome-wide RNAi screens in cultured fly cells identified just 18 proteins that, when depleted, gave a reduced number of centrioles (Goshima et al., 2007; Dobbeltrae et al., 2008). This list includes Sak, DSas-6, and DSas-4, as well as eight other proteins that specifically localize to centrosomes (Ana1, Ana2, Ana3, Asl, DCP110, DCP135/Bl10, DCEP97, and Rcd4); these eight are therefore good candidates to play a direct role in centriole duplication.

GFP-Sak, GFP–DSas-6, and DSas–4–GFP share the unique ability to drive de novo formation of centriole-like structures in unfertilized eggs when highly overexpressed from the upstream activation sequence (UAS) promoter (Peel et al., 2007; Rodrigues-Martins et al., 2007b). UAS-GFP-Sak and UAS-GFP–DSas-6 induce these structures in \( \sim \)95% of unfertilized eggs, whereas UAS–DSas–4–GFP does so in \( \sim \)60% of unfertilized eggs (Peel et al., 2007). We wondered if we could use this assay to identify other components likely to function upstream in the centriole duplication pathway. We therefore generated transgenic lines carrying GFP fusions to all eight potential duplication factors under the control of the UAS promoter, which allowed us to overexpress them in unfertilized eggs (Fig. S1). Strikingly, only Ana2 (in 97% of eggs) and Asl (in 33% of eggs) were able to drive de novo formation of centriole-like structures (Fig. 1).

Asl has recently been shown to be essential for centriole duplication in flies (Blachon et al., 2008), whereas, of the six proteins unable to induce de novo centriole formation, two, DCEP135/Bl10 and Ana3, are now known not to be essential for centriole duplication in flies (Mottier-Pavie and Megraw, 2009; Stevens et al., 2009). These findings indicate that our overexpression assay can identify those proteins likely to be most intimately involved in centriole duplication. As Asl has already been shown to be required for centriole duplication (Blachon et al., 2008), we focused on investigating the function of Ana2.

**Ana2 is an important regulator of canonical centriole duplication**

Ana2 can drive de novo formation of centriole-like structures as efficiently as DSas-6 and Sak (Peel et al., 2007; Rodrigues-Martins et al., 2007b). We wanted to verify, however, that it also has a role in canonical centriole duplication. Overexpressing GFP-Sak or GFP–DSas-6 from the ubiquitin (Ubq) promoter induces centriole overduplication in brains and embryos, respectively (Peel et al., 2007). Surprisingly, however, overexpression of Sak, DSas-6, or DSas-4 cannot drive centriole overduplication in primary spermatocytes (Peel et al., 2007), which suggests that another duplication protein is limiting. To test if Ana2 might be this limiting factor, we generated Ubq-GFP-Ana2 transgenic lines. Strikingly, we found that in spermatocytes expressing Ubq-GFP-Ana2, in addition to the normal centriole pairs (doublets), we observed centriole triplets, quadruplets, and even quintets (Figs. 2, A–G). The extra centrioles in these clusters appeared to be fully functional; they separated from one another by the end of meiosis I (as centriole doublets normally do), and the extra centrioles inherited by secondary spermatocytes recruited PCM and nucleated MT asters, and so formed multipolar spindles during meiosis II (Fig. 2, H and I).

We did not observe centriole overduplication in embryos or brain cells expressing Ubq-GFP-Ana2 (unpublished data), which is consistent with DSas-6 and Sak levels, respectively, limiting centriole formation in these tissues. Nevertheless, that Ana2 overexpression can drive centriole overduplication in spermatocytes demonstrates that it is an important regulator of canonical centriole duplication.

**Ana2 shows a unique asymmetric localization to the daughter centriole**

We next wanted to compare the localization of Ana2 with that of the other *Drosophila* centriole duplication factors. DSas–4–GFP, GFP–DSas-6, and GFP-Sak are all enriched at the proximal and distal ends of the large spermatocyte centrioles (Peel et al., 2007). We found that, likewise, Ana2-GFP localized preferentially to the proximal and distal centriole tips. Strikingly, however, Ana2-GFP (and GFP-Ana2) also exhibited a unique asymmetric distribution, consistently localizing preferentially along one centriole barrel (Fig. 3, A and B).

In primary spermatocytes, it is possible to distinguish mother and daughter centrioles, as the daughter can often be observed associating end-on with the side of the mother (Blachon et al., 2008). In 25 centriole pairs where we could unambiguously distinguish mother and daughter centrioles, Ana2-GFP was always enriched on the daughter (Fig. 3 A). Mother and daughter centrioles can show important differences in their behavior in vertebrate cells (Piel et al., 2000) and during asymmetric stem cell divisions in *Drosophila* (Reblolo et al., 2007; Rusan and Peifer, 2007; Yamashita et al., 2007). Although mother and daughter centrioles are morphologically and molecularly distinguishable in vertebrates (see, for example, Vorobjev and Chentsov, 1982; Chang et al., 2003; Gromley et al., 2003; Graser et al., 2007), this is not the case in *Drosophila* (Callaini and Riparbelli, 1990; Callaini et al., 1997; Vidwans et al., 2003). To our knowledge, Ana2-GFP is the first fly protein shown to localize asymmetrically to mother and daughter centrioles in this manner.

Interestingly, as spermatocytes progressed through meiosis I, this centriolar asymmetry became less pronounced, and this appeared to reflect the selective loss of GFP-Ana2 from the daughter centriole, bringing its levels down to that of the mother (compare Fig. 3 B, showing a G2 centriole pair, to Fig. 3 C,
showing a centriole pair separating at the end of meiosis I). As overexpression of Ana2 can lead to centriole overduplication, Ana2 levels presumably must normally be tightly regulated to prevent the formation of extra centrioles.

After exit from meiosis II, each spermatid inherits a single centriole, which acts as a basal body to nucleate the flagellar axoneme. Structural components of the centriole, like Ana3 (Stevens et al., 2009) and *Drosophila* pericentrin-like protein (D-PLP; Martinez-Campos et al., 2004), continue to localize along the basal body. In contrast, Ana2, like the conserved duplication proteins (Blachon et al., 2009), was undetectable along the basal body (Fig. 3 D). Ana2 did, however, colocalize with GFP–DSas-6 at the proximal centriole-like structure (Fig. 3 D), a small nodule adjacent to the basal body that has been proposed to be an early intermediate in centriole formation (Blachon et al., 2009).
Ana2 is the likely functional orthologue of <i>Caenorhabditis elegans</i> SAS-5

Intriguingly, <i>Drosophila</i> homologues have been identified for all the <i>C. elegans</i> centriole duplication factors except SAS-5, which has no clear homologues outside worms. Ana2 and SAS-5 are similar in size and have a single central coiled-coil domain, leading Goshima et al. (2007) to suggest that Ana2 could be the <i>Drosophila</i> equivalent of SAS-5 (Goshima et al., 2007). As SAS-5 interacts with SAS-6 in worms (Leidel et al., 2005), we looked for a genetic interaction between Ana2 and DSas-6 in flies.

Figure 2. Overexpression of Ana2 drives centriole overduplication in spermatocytes. (A and B) Centriole number (A) and conformation (B) in G2 primary spermatocytes expressing either the centriole marker RFP-PACT alone or both RFP-PACT and GFP-Ana2. Centrioles were counted in a total of 109 RFP-PACT cells and 138 GFP-Ana2 RFP-PACT cells from seven testes per condition. (C and D) G2 primary spermatocytes expressing either RFP-PACT (red) alone (C) or both RFP-PACT and GFP-Ana2 (D). DNA is in blue. The cell in C has the normal two centriole pairs. Overexpression of GFP-Ana2 induces centriole triplets and quadruplets (D). (E–G) Magnified images of RFP-PACT–labeled doublet (E), triplet (F), and quadruplet (G) centriole groups. (H and I) Secondary spermatocytes in meiosis II expressing either RFP-PACT (red) alone (H) or both RFP-PACT and GFP-Ana2 (I). Tubulin is in green and DNA in blue. The cell in H has the normal two centrioles whereas the one in I has three centrioles forming a tripolar spindle. Bars: (C and D) 10 µm; (H and I) 5 µm.
Interestingly, the centriole-like structures produced by overexpressing UASp-GFP–DSas-6 differ significantly from those resulting from the overexpression of GFP-Sak, DSas-4–GFP, Asl-GFP, or Ana2-GFP in that they are much larger and often appear ring-shaped, and that only one structure is contained within each aster (Fig. 4 E; Peel et al., 2007; Rodrigues-Martins et al., 2007a). The structures in the eggs from females expressing both Ubq-GFP–DSas-6 and Ubq-Ana2-GFP were similar to this DSas-6 type (Fig. 4, D and E). Importantly this interaction was specific to Ana2 and DSas-6. In eggs from mothers carrying one copy of either Ubq-Ana2-GFP or Ubq-GFP–DSas-6 together with one copy of either Ubq-GFP-Sak, Ubq-Asl-GFP, or Ubq–DSas-4–GFP, we observed at most a very small number of asters in very few eggs (Fig. 4 A).

A small percentage of eggs laid by mothers carrying two copies of a Ubq-GFP–DSas-6 transgene (as opposed to the much stronger UASp-GFP–DSas-6 discussed above) assemble centriole-like structures (Peel et al., 2007). To see if we could enhance this effect, we generated flies carrying one copy of Ubq-GFP–DSas-6 and one copy of Ubq-Ana2-GFP, neither of which alone (as a single copy) induces the assembly of centriole-like structures (Fig. 4 A). Strikingly, almost all the unfertilized eggs laid by these females contained hundreds of large structures that stained for centriole markers, recruited PCM, and nucleated asters (Fig. 4, A–D; and Fig. S2 A). Importantly, this interaction was specific to Ana2 and DSas-6. In eggs from mothers carrying one copy of either Ubq-Ana2-GFP or Ubq-GFP–DSas-6 together with one copy of either Ubq-GFP-Sak, Ubq-Asl-GFP, or Ubq–DSas-4–GFP, we observed at most a very small number of asters in very few eggs (Fig. 4 A).

Figure 3. Ana2 is a centriole component with a unique asymmetric localization. (A) Centriole pair from a G2 primary spermatocyte expressing Ana2-GFP (green) stained for the centriole marker GTU88* (red). Ana2-GFP localizes to the proximal and distal centriole ends and also exhibits a unique asymmetric distribution, localizing preferentially along one centriole barrel, which can be identified as the daughter from the GTU88* staining (see main text). (B) Centriole pair from a G2 primary spermatocyte expressing GFP-Ana2 (green) and RFP-PACT (red). GFP-Ana2 localization is indistinguishable from Ana2-GFP. (C) Centriole pair from a primary spermatocyte at anaphase of meiosis I: the centrioles are beginning to separate. The cell is expressing RFP-PACT (red) and GFP-Ana2 (green), which is no longer obviously asymmetric. (D) Two basal bodies from spermatids expressing RFP-PACT (blue) and GFP–DSas-6 (green), and stained for Ana2 (red). GFP–DSas-6 and Ana2 colocalize at the proximal centriole-like structure, a nodule adjacent to the basal body marked by RFP-PACT. Bars, 2 µm.
Figure 4. Ana2 and DSas-6 functionally and physically interact. (A) Percentage of unfertilized eggs laid by mothers of the given genotypes that contained MT asters. All transgenes were GFP fusions with a Ubq promoter. Eggs from mothers expressing one or two copies of Ubq-GFP–DSas-6 and Ubq-Ana2-GFP were analyzed; all combinations expressed one copy of each transgene. n > 80 eggs per genotype (for values, see Materials and methods). (B and C) Almost all unfertilized eggs from mothers expressing one copy of Ubq-GFP–DSas-6 and one copy of Ubq-Ana2-GFP assemble large numbers of
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Ana2 is the most similar Drosophila protein. Although vertebrate STIL family members are larger than Ana2 or SAS-5, all of these proteins share a short, central, coiled-coil domain (Fig. 5). In addition, we identified a particularly conserved region of \( \approx 90 \) aa toward the C terminus of Ana2 and STIL, which we called the STAN motif. The STAN motif of Ana2 is 31% identical (48% similar) to that of zebrafish STIL. A divergent STAN motif can be detected in SAS-5, which is 12% identical (26% similar) to that of zebrafish STIL (Fig. 5). Importantly, the STAN motif is within the regions of SAS-5 and Ana2 that interact with SAS-6 and DSas-6, respectively (Fig. 4 F; Boxem et al., 2008).

Data from studies of STIL in mice, zebrafish, and humans are consistent with a function in centriole duplication, although this was not appreciated at the time of these studies. First, mitotic spindles often lack centrosomes in stil mutant zebrafish (Pfaff et al., 2007). Second, STIL mutant mice show defects characteristic of aberrant cilia function, such as randomized left–right asymmetry and neural tube abnormalities.

Figure 5. Ana2 is related to vertebrate STIL. Schematic of human STIL Drosophila Ana2, and C. elegans SAS-5. All three proteins have a central, coiled-coil domain (green) and a conserved region near the C terminus (blue): the STAN motif. An alignment of the STAN motif is shown in full, with an alignment including SAS-5 below. Both are colored according to the Blosum62 coloring scheme, where dark blue indicates a match to the consensus sequence and light blue indicates a positive Blosum62 score. Asterisks indicate residues are identical in all aligned sequences, colors indicate conserved substitutions, and periods indicate semiconserved substitutions.

interaction (Fig. 4 F and Fig. S2 B). Moreover, like SAS-5, Ana2 also interacts with itself. Our attempts to test whether Ana2 and DSas-6 associate in vivo were hindered by their low abundance. However, we found that DSas-6 antibodies communoprecipitated Ana2-GFP from S2 cells overexpressing Ana2-GFP (Fig. 4 G). Collectively, our evidence of a specific functional and physical interaction between Ana2 and DSas-6 indicates that Ana2 likely represents the Drosophila functional orthologue of SAS-5.

Ana2 is related to the human protein SCL/TAL1 interrupting locus (STIL/SIL), which is mutated in primary microcephaly. Having shown that Ana2 is the likely SAS-5 functional orthologue in Drosophila, we looked for Ana2/SAS-5 orthologues in other species. Using an iterative basic local alignment search tool (BLAST) search, we found significant homology between Ana2 and the STIL or SIL protein family. Moreover, the reciprocal iterative BLAST search starting with zebrafish STIL identified Ana2 as the most similar Drosophila protein. Although vertebrate STIL family members are larger than Ana2 or SAS-5, all of these proteins share a short, central, coiled-coil domain (Fig. 5). In addition, we identified a particularly conserved region of \( \approx 90 \) aa toward the C terminus of Ana2 and STIL, which we called the STil/ANa2 (STAN) motif. The STAN motif of Ana2 is 31% identical (48% similar) to that of zebrafish STIL. A divergent STAN motif can be detected in SAS-5, which is 12% identical (26% similar) to that of zebrafish STIL (Fig. 5). Importantly, the STAN motif is within the regions of SAS-5 and Ana2 that interact with SAS-6 and DSas-6, respectively (Fig. 4 F; Boxem et al., 2008).

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Apart how these components cooperate to build a centriole of known to cause MCPH, and all are centrosomal proteins (Bond genes, produced brain size (Kumar et al., 2009). Mutations in four other genes, MCPH1, CDK5RAP2, ASPM, and CPAP/CENPJ, are known to cause MCPH, and all are centrosomal proteins (Bond et al., 2002, 2005; Jackson et al., 2002; Zhong et al., 2005, 2006), which strongly suggests that STIL is required for efficient centrosome function in humans.

Conclusions

Here, we show that of eight centrosomal proteins identified as potential duplication factors in Drosophila tissue culture cells, only two, Asl and Ana2, appear to be able to induce de novo formation of centriole-like structures in unfertilized eggs. Asl has recently been shown to be essential for centriole duplication (Blachon et al., 2009), and we provide evidence that Ana2 is also a key centriole duplication factor. Thus, Ana2 and Asl join Sak, DSas-6, and DSas-4 to make up a module of just five proteins known to drive centriole duplication in flies.

Our data strongly suggest that the Ana2/STIL family of centrosomal proteins are the long-sought functional orthologues of SAS-5. Thus, four of these five components (Sak/ZYG-1, DSas-6/SAS-6, Ana2/SAS-5, and DSas-4/SAS-4) are functionally conserved between flies and worms. Moreover, three of these proteins are required for centriole duplication in humans (Habedanck et al., 2005; Leidel et al., 2005; Kleylein-Sohn et al., 2007; Smit et al., 2007; Kohlmaier et al., 2009), whereas the fourth, SAS-5/Ana2/STIL, also appears likely to be required for this process in vertebrates (Izraeli et al., 1999; Pfaff et al., 2007; Kumar et al., 2009).

Both flies and worms have an additional protein (SPD-2 in worms, Asl in flies) that appears to be essential for centriole duplication. Intriguingly, both SPD-2 (Kemp et al., 2004; Pelletier et al., 2004) and Asl (Bonaccorsi et al., 1998; Varmark et al., 2007; Blachon et al., 2008) are not only required for centriole duplication, but also for PCM recruitment. There is evidence that the PCM promotes centriole duplication (Dammersmann et al., 2004; Loncarek et al., 2008), so SPD-2 and Asl could play a more indirect role in centriole duplication via their ability to recruit PCM. Alternatively, both proteins may act directly in centriole duplication, with the function of SPD-2 in worms perhaps being performed by Asl in flies.

In summary, we show that Ana2 acts as a centriole duplication factor in Drosophila and is likely to have a conserved role in other species. Overall, centriole duplication appears to be a highly conserved process, at the heart of which is a small number of key proteins. The challenge will now be to tease apart how these components cooperate to build a centriole of the right size, in the right place, and at the right time.

Materials and methods

Generation of GFP fusions and transgenic lines

P element-mediated transformation vectors containing GFP-fusions to Ana1, Ana2, Ana3, Asl, DCP110, DCEp135, DCEp97, and Rcd4 were generated as follows. The complete coding region of each protein was amplified from either cDNA (Ana1, Ana2, Asl, DCP110, DCEp135, and Rcd4) or genomic DNA (Ana3 and DCEp97), with att sites at either end for Gateway cloning (Invitrogen). These fragments were inserted into the Gateway pDONR Zero vector. The two Ana3 exons were amplified separately from genomic DNA before being ligated together to produce a vector with the complete coding sequence. The pDONR vectors were then recombined with UAosp and, for Ana2 and Asl, Ubq plasmids (Peel et al., 2007), with each coding sequence placed in frame with GFP at the N or C terminus. The following transgenic lines were generated by BestGene, Inc.: UAosp-GFP-Ana1, UAosp-Ana2-GFP, UAosp-Ana3-GFP, UAosp-Asl-GFP, UAosp-DCP110-GFP, UAosp-GFP-DCEp135, UAosp-GFP-DCEp97, UAosp-GFP-Rcd4, Ubq-Ana2-GFP, Ubq-GFP-Ana2, and Ubq-Asl-GFP. The Ubq promoter drives moderate expression in all tissues (Lee et al., 1988), whereas the UAosp lines were crossed to V32a, which expresses a Gal4/VP16 fusion protein from a maternal tubulin promoter; this drive a very high-level over-expression in the female germline (Peel et al., 2007).

We also used the previously described transgenic lines Ubq-GFP–DSas-6, Ubq-GFP-Sak, Ubq-Dasas-4-GFP, and Ubq-MRP-pericentriolar/AKAP450 centrosomal-targeting domain (PACT; Peel et al., 2007).

Generation and use of Ana2 antibodies

A maltose-binding protein (MBP, New England Biolabs, Inc.) fusion of an 1–201 of Ana2 was purified according to the manufacturer’s instruction, and antisera were raised in two rabbits by Eurogentec. To affinity purify antibodies, the antisera was first depleted of anti-MBP antibodies by passing over a ProteinA AffiGel MBP column (Thermo Fisher Scientific). Specific antibodies were then purified by passing the antisera over a column of MBP-Ana2(1–201) fusion protein. The column was washed with PBS + 0.5 M NaCl, and antibodies were eluted in 0.1 M glycine, pH 2.1. The antibodies were neutralized with 1 M Tris, pH 8.5, and glycerol was added to 50%, then materials were stored at −20°C.

The antibody was used at 1:250 for immunofluorescence experiments. It weakly stained centrosomes in embryos and the proximal centriole-like structure in spermatozoon, but it did not stain primary spermatocyte centrosomes. To investigate why this was the case, we used the antibody to stain Ana2-GFP-expressing spermatids. Here, the antibody stained the distal tips of the centrioles, but not the proximal ends of the centrioles or the single centriole barrel, even though the Ana2-GFP labeling was clearly visible at these sites. This suggests that the antibody does not stain spermatocyte centrosomes for a combination of reasons. First, endogenous Ana2 must be present at centrioles at very low levels, as we cannot detect it even in the distal portion of the centrioles where we can detect Ana2-GFP. Second, endogenous Ana2 is probably not easily accessible to antibodies at the proximal end of the centrioles and along the centriole barrel, as we cannot detect Ana2-GFP with the antibody at these sites even though Ana2-GFP is localized there. For other uses see the “Electrophoresis and immunoblotting” and “Immunoprecipitation” sections.

Fixed analysis of eggs and pupal testes

0–4 h of collection of unfertilized eggs were made from mothers expressing UAosp-GFP-Ana1 (n = 77), UAosp-Ana2-GFP (n = 123), UAosp-Ana3-GFP (n = 206), UAosp-Asl-GFP (n = 165), UAosp-DCP110-GFP (n = 239), UAosp-GFP-DCEp135 (n = 177), UAosp-GFP-DCEp97 (n = 172), UAosp-GFP-Rcd4 (n = 214), Ubq-GFP-DSas-6 (n = 90), Ubq-GFP-DSas-6/2 copies (n = 92), Ubq-Ana2-GFP (n = 90), Ubq-Ana2-GFP 2 copies (n = 84), Ubq-GFP–DSas-6/Ubq-Ana2-GFP (n = 84), Ubq-GFP–DSas-6/Ubq-Dasas-4–GFP (n = 81), Ubq-GFP–DSas-6/Ubq-GFP-Sak (n = 81), Ubq-GFP–DSas-6/Ubq-Asl-GFP (n = 96), Ubq-Ana2-GFP/Ubq-Dasas-4-GFP (n = 117), Ubq-Ana2-GFP/Ubq-GFP-Sak (n = 96), and Ubq-Ana2-GFP/Ubq-Asl-GFP (n = 86). Eggs were dechorionated in 60% bleach for 2 min, washed in water + 0.05% Triton X-100, then washed into a small glass bottle with 1 ml heptane. 1 ml methanol + 5% 0.25 M EGTA was added and the bottle was shaken gently until most eggs fell into the lower methanol/EGTA layer. Eggs were stored in methanol at 4°C. For immunostaining, eggs were rehydrated by washing in PBT (PBS + 0.1% Triton X-100), blocked in PBS + 5% BSA, and incubated with primary antibodies in PBS/BSA overnight at 4°C. Eggs were then washed in PBT before incubation with secondary antibodies diluted 1:1,000 in PBT for 4 h at room temperature. After final washes in PBT, eggs were mounted in mounting medium (85% glycerol and 2.5% propyglycollate).

Pupal testes were dissected in PBT, placed on a coverslip, and cut open. A slide was then placed over the coverslip and then flash frozen in liquid nitrogen. Coverslips were removed and the slides were incubated for 5 min in methanol at −20°C, and then in acetone for 1–2 min at −20°C. This was followed by incubation in PBT for 10 min, washes in PBS, and blocking in PBS/BSA (1%). Slides were incubated in primary antibody
An 80-cm³ flask of cells was grown for 3 d and harvested by centrifugation (1 h at 25°C). After final washes, slides were mounted in mounting medium.

Slides were observed at room temperature on a spinning disc confocal system [ERS; PerkinElmer], mounted on an inverted microscope (Axiovert 200M; Carl Zeiss, Inc.) with a charge-coupled device camera (Orca, Hamamatsu), using a 63×/1.2 NA objective (Carl Zeiss, Inc.) and Immersol oil (Carl Zeiss, Inc.). Images were acquired using Ultraview ERS software (PerkinElmer), imported into Photoshop CS2 (Adobe), and adjusted to use the full range of pixel intensities.

Identification of centriole-like structures in eggs

Unfertilized eggs were initially stained for α-tubulin and the centriole marker DSas-4. In wild-type (WT) eggs, the only MTs visible are those surrounding the polar bodies (Peel et al., 2007; Stevens et al., 2009). For these conditions, we went on to stain unfertilized eggs with antibodies against a second centriole marker, DPF, and the PCM proteins centrosomin [Cnn] and γ-tubulin. Structures were considered to be centriole-like if they stained for both of the centriole and both of the PCM markers, and nucleated MT asters. We then quantified the percentage of unfertilized eggs containing these structures for each overexpression condition.

Antibodies

The following antibodies were used: 1:1,000 rabbit anti–DPF (Martinez-Campos et al., 2004), 1:250 rabbit anti–DSas-4 (Basto et al., 2006), 1:500 guinea pig anti–Cnn (Dix and Raff, 2007), 1:100 mouse monoclonal anti–α-tubulin (DM1a; Sigma-Aldrich), and 1:1,000 GTUB88*, a batch of the mouse monoclonal anti–γ-tubulin GTUB88 (Sigma-Aldrich) antibody that cross-reacts with centrioles in flies (Martinez-Campos et al., 2004); Alexa Fluor 488, Cy3, and Cy5 secondary antibodies were obtained from Invitrogen or Jackson ImmunoResearch Laboratories, Inc.

Y2H assay

pDEST22 [prey] and pDEST32 [bait] vectors containing full-length Ana2, full-length DSas-6, DSas-6 NT (aa 1–210), DSas-6 M (aa 104–317), DSas-6 CT (aa 224–414), Ana2, Ana2 M (aa 201–414, 1× PMSF, 1× protease inhibitor [Roche], and 1% penicillin/streptomycin (Invitrogen). pUbq-Ana2-GFP vectors were transfected into S2 cells using Cellfectin (Invitrogen) using the pCoBlast vector for selection. After 3 wk of selection with blasticidin, stable cell lines were obtained. The membrane was washed in PBS and incubated with horseradish peroxidase-conjugated secondary antibodies (GE Healthcare) diluted 1:1,500,000 in PBS for 1 h at room temperature. Finally, the membrane was washed in PBS, incubated with ECL advance chemiluminescent substrate (GE Healthcare) according to manufacturer’s instructions, and exposed to x-ray film.

Identification of Ana2 homologues and sequence alignments

The position-specific iterated BLAST (PSI-BLAST) algorithm (Altschul et al., 1997) from the National Center for Biotechnology Information was used to search for homologues of Ana2. Multiple sequence alignments were performed using ClustalW2 (Larkin et al., 2007) and visualized in Jalview (Waterhouse et al., 2009) using the Blossum2 coloring scheme.

Online supplemental material

Fig. S1 is a Western blot showing that both UASp-GFP-Ana1 and UASp-Ana2-GFP are overexpressed at very high levels in unfertilized eggs. Fig. S2 shows the raw data from our Y2H analysis of Ana2 and DSas-6. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200910016/DC1.

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References


a major determinant of cerebral cortical size. Nat. Genet. 32:316–320. doi:10.1038/ng995


Figure S1. Overexpression of centrosomal proteins in eggs. The UAS-Gal4 system can produce extremely high levels of overexpression in eggs. Western blots of eggs laid by WT, UAS-GFP-Ana1, and UAS-Ana2-GFP mothers are shown as an example. Serial dilutions of the UAS-GFP-Ana1 and UAS-Ana2-GFP egg extracts show that both proteins are overexpressed by a factor of >100. Neither endogenous Ana1 nor Ana2 is detectable. Actin is shown as a loading control.
Figure S2. **Ana2 and DSas-6 interact functionally to induce de novo formation of centriole-like structures and physically by a Y2H assay.** (A) Single aster from an egg laid by a Ubq-GFP–DSas-6/Ubq-Ana2-GFP mother stained for tubulin (blue), Cnn (green), and D-PLP (red). The aster surrounds a ring-shaped structure that stains for the centriole marker D-PLP and recruits the PCM protein Cnn. Bar, 2 µm. (B) Y2H assay testing full-length (FL) Ana2 and DSas-6 as well as N-terminal (NT), C-terminal (CT), and middle (M) fragments of DSas-6, and NT and CT fragments of Ana2. Rows 1, 2, 3, and 4 are as follows: 1, selection on −Leu, −Trp, −His with 5 mM 3AT; 2, selection on −Leu, −Trp, −His with 20 mM 3AT; 3, selection on −Leu, −Trp, −Ade; and 4, X-gal assay. Asterisks indicate positive results due to autoactivation.