The immunological synapse (IS) defines the interface formed between an immune cell and the cell with which it is interacting. It was originally defined in T cells but has subsequently been characterized in many other types of immune cells. There are several proposed functions for the IS that are likely to vary depending on the cell type. These range from signal generation to termination of effector function, but their relative importance has been debated (1). One of the clearest and first appreciated functions of the IS, however, is directed secretion (2, 3). Directed secretion enables precise control in elaborating cell contents onto a specific target cell. This may be especially useful if the secreted components have the potential to harm cells, as in cytotoxicity. In this case, directed secretion serves the purpose of delivering lytic molecules to a target cell while protecting neighboring cells. Directed secretion requires numerous cell surface and intracellular alterations that facilitate the movement of secretory components to the IS. The polarization of secretory machinery in cytotoxic cells, such as NK cells, toward a target cell has long been appreciated (4). The process of directed secretion for cytotoxicity culminates with the release of lytic granules, which are secretory lysosomes containing perforin and granzymes (5) onto the target cell. This is required for host defense, as at least three human genetic defects that block this process (mutations in the genes encoding Rab27a, Munc13-4, and syntaxin-11) result in impaired cytotoxicity and susceptibility to viral infection (6–8).

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### Cdc42-interacting protein–4 functionally links actin and microtubule networks at the cytolytic NK cell immunological synapse

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An essential function of the immunological synapse (IS) is directed secretion. NK cells are especially adept at this activity, as they direct lytic granules to the synapse for secretion, which enables cytotoxicity and facilitates host defense. This initially requires rearrangement of the actin cytoskeleton and, subsequently, microtubule-dependent trafficking of the lytic granules. As these two steps are sequential, specific linkages between them are likely to serve as critical regulators of cytotoxicity. We studied Cdc42-interacting protein–4 (CIP4), which constitutively interacts with tubulin and microtubules but focuses to the microtubule organizing center (MTOC) after NK cell activation, when it is able to associate with Wiskott-Aldrich syndrome protein (WASp) and the actin filament–rich IS. WASp deficiency, overexpression of CIP4, or parts of CIP4 interfere with this union and block normal CIP4 localization, MTOC polarization to the IS, and cytotoxicity. Reduction of endogenous CIP4 expression using small interfering RNA similarly inhibits MTOC polarization and cytotoxic activity but does not impair actin filament accumulation at the IS, or Cdc42 activation. Thus, CIP4 is an important cytoskeletal adaptor that functions after filamentous actin accumulation and Cdc42 activation to enable MTOC polarization and NK cell cytotoxicity.
lytic granules to the IS, however, is a relatively late step in the formation of the IS and requires several preceding steps, including reorganization of the actin cytoskeleton, clustering of cell-surface receptors, and polarization of the microtubule organizing center (MTOC). Movement of the MTOC to the IS in CTLs is essential for lytic granule delivery and release (11), and NK cells in which the MTOC does not polarize are unable to mediate cytotoxicity (12, 13). In T cells, polarization of the MTOC requires the action of the Cdc42 GTPase (14), and in NK cells it is dependent on activation-induced signaling (13). Although Cdc42 activity is also required for normal filamentous actin (F-actin) accumulation at the IS (14, 15), microtubule integrity is not. In NK cells, the F-actin–rich supramolecular activation cluster (SMAC) forms normally after microtubule depolymerization, but SMAC formation and lytic granule polarization are both blocked by inhibition of actin function (10). This suggests that F-actin reorganization in the SMAC is an early and upstream step in the process of NK cell IS formation and function. Similarly, NK cells from patients with Wiskott-Aldrich syndrome (WAS) who lack the WAS protein (WASp), which is required for F-actin reorganization, have reduced F-actin in the SMAC and decreased polarization of lytic granules (16). Although WASp–dependent F-actin reorganization is an essential prerequisite, reorientation of the MTOC to the SMAC is a likely critical late step in cytotoxicity. In particular, the ability to link these two steps involving the reorganized actin cytoskeleton and microtubule network in NK cells could help facilitate the directed secretion of lytic granules. A specific protein with these capabilities in cytotoxic cells, however, has not been demonstrated, but would define an important regulatory step in formation of the cytolytic IS and cytotoxicity.

We have studied Cdc42–interacting protein–4 (CIP4) as a potential candidate to link linear stages in cytolytic–directed secretion. CIP4 was discovered for its ability to interact with Cdc42 and with WASp through an Src homology 3 (SH3) domain (17, 18). It also contains a Fes/CIP4 homology (FCH) domain, which enables it to interact with tubulin, and may be relevant to podosome formation in macrophages (19). In this study, we have determined that appropriate expression of CIP4 in NK cells is required for MTOC polarization and cytotoxicity. We show that CIP4 biochemically and spatially unites F-actin and microtubule networks in activated NK cells but functions downstream of Cdc42 activation. Thus, we propose that CIP4 is an important functional linkage between the SMAC and MTOC polarization to obtain secretory cytolytic effector function.

RESULTS

CIP4 is expressed in NK cells and localizes with the MTOC to the IS after NK cell activation

To determine if CIP4 could serve a role in NK cells, we initially documented its expression. Using overlapping primer sets that encompassed the complete coding sequence, a CIP4 message was identified among cDNA in all immortalized NK cell lines tested, as well as in ex vivo NK cells (Fig. 1 A). Sequence analysis of amplified product confirmed consensus in all NK cell samples. CIP4 was also detected by Western blotting among total lysates using mAb anti-CIP4 in NK cell lines and ex vivo NK cells, albeit at levels lower than those found in fibroblasts (not depicted) (Fig. 1 B). Levels of WASp were more similar among the different NK cells and, unlike CIP4, were proportional to α-tubulin, which is shown as a protein loading control. To more accurately determine the relative levels of CIP4 in NK cells, intracellular FACS for CIP4 was developed (Fig. 1 C). Relative to isotype control, CIP4 content was uniform in the different NK cell lines but was lower in NK cells among ex vivo PBMCs (Fig. 1 D). This result was consistent with the smaller size of ex vivo NK cells compared with the NK cell lines (unpublished data).

CIP4 has important associations with WASp and microtubules in nonimmune cells, as well as in macrophage podosomes (18, 19). Because we have identified critical roles for WASp and microtubules in the polarization of lytic granules to the IS in NK cells (10, 16), we wanted to study CIP4 relative to the IS. As an initial approach, we used YTS NK cells stably expressing a CD2-GFP fusion molecule, as CD2 clustering serves as an early marker of the IS in NK cells (10). Like certain subpopulations of ex vivo NK cells, YTS cells can be induced to mediate cytotoxicity after ligation of CD28 (20, 21). Because K562 cells do not express CD80 or CD86, they are not susceptible to lysis by YTS cells (unpublished data). To use K562 cells as a susceptible target for YTS cells that would induce a cytolytic IS, the K562 cells were stably transduced with CD86 (KT86 cell line). KT86 cells were killed by YTS cells, as measured in 51Cr release assays, and provide the advantage of isolating a specific activation pathway in YTS cells. Direct comparison of KT86 and K562 cells also affords a comparison of cytolytic and noncytolytic interactions.

Using YTS CD2-GFP cells, the localization of CIP4 relative to the IS and to the MTOC was determined by confocal microscopy in cytolytic (Fig. 2, A–F) and noncytolytic (Fig. 2, G–L) conjugates. In this setting, the MTOC was visualized as an accumulation of α-tubulin and was defined using an intensity and size threshold, or as the region containing the centrosomal protein pericentrin (Fig. 2, C and I). In cytolytic conjugates formed between YTS and KT86 cells, the IS as defined by the presence of accumulated CD2-GFP contained a polarized tubulin–defined MTOC, and accumulated CIP4 (Fig. 2, A–F). The noncytolytic synapse also contained accumulated CD2–GFP but had reduced MTOC polarization and accumulation of CIP4 (Fig. 2, G–L). In experiments where the nucleus was visualized, the MTOC and CIP4 could be found between the nucleus and the IS in YTS CD2-GFP cells engaged in cytolytic, but not in noncytolytic, conjugates (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20061893/DC1). Assessment of multiple conjugates over separate experiments demonstrated that although CD2-GFP accumulation at the IS was slightly decreased in noncytolytic compared with cytolytic synapses (by 14%), polarization of
between /H9251-tubulin and the tubulin-defined MTOC were not significant (P = 0.22), further analyses focused on the tubulin-defined MTOC. Colocalization analyses were also performed using a pericentrin-defined MTOC. There was 75 – 97% colocalization between pericentrin and the tubulin-defined MTOC but only 45% colocalization between the tubulin-defined MTOC and the pericentrin-defined MTOC (Fig. 2 Q). Thus, the area occupied by the pericentrin-defined MTOC was less than the tubulin-defined MTOC. Colocalization between the pericentrin-defined MTOC and CIP4, however, was greater in cytolytic conjugates than the colocalization between CIP4 and the tubulin-defined MTOC. This difference was not found in noncytolytic conjugates, as CIP4 colocalization with the MTOC was less (note MTOC enlargement in Fig. S2, available at http://www.jem.org/cgi/content/full/jem.20061893/DC1) and, thus, is consistent with the centrosomal focusing of CIP4 in cytolytic conjugates. As an alternative to these whole-cell area and intensity cell-based colocalization analyses, the colocalization within a constant size region around the MTOC was also performed and defined a similarly increased colocalization between the MTOC and CIP4 in YTS CD2-GFP cells engaged in cytolytic versus noncytolytic conjugates (Fig. S2).

Because CIP4 could have the potential to facilitate interaction between the MTOC and F-actin during formation of the MTOC and accumulation of CIP4 were substantially reduced (by 72%; Fig. 2 M). To assess MTOC polarization in this system more definitively, the mean distance of the tubulin- or pericentrin-defined MTOC from the IS was also measured in cytolytic and noncytolytic conjugates. The MTOC was significantly closer to the IS in cytolytic conjugates (Fig. 2 N), consistent with the observed increase in MTOC polarization and with the role of the MTOC in facilitating lytic granule delivery (11).

To quantitatively assess colocalization between CIP4 and the MTOC, the area occupied by the different fluorescent intensities within the YTS cells was measured. The area of the tubulin-defined MTOC was distinguished from the fluorescence of unaccumulated α-tubulin and individual microtubules, but in the plane of the MTOC contains the majority of the α-tubulin fluorescent area (Fig. 2 O). The area occupied by α-tubulin, the tubulin-defined MTOC, or CIP4 did not differ between YTS cells in cytolytic and noncytolytic conjugates (Fig. 2 O). In contrast, the colocalization between α-tubulin, or the tubulin-defined MTOC, and CIP4 was increased in cytolytic conjugates (Fig. 2 P). Similarly, the colocalization between CIP4 and the tubulin-defined MTOC or α-tubulin was also higher in cytolytic conjugates. Because the area of CIP4 did not increase in cytolytic YTS cells, this suggested that the existing CIP4 focused to the MTOC during the cytolytic process. As differences in CIP4 colocalization between α-tubulin and the tubulin-defined MTOC were not significant (P = 0.22), further analyses focused on the tubulin-defined MTOC.

Colocalization analyses were also performed using a pericentrin-defined MTOC. There was 75 – 97% colocalization between pericentrin and the tubulin-defined MTOC but only 45% colocalization between the tubulin-defined MTOC and the pericentrin-defined MTOC (Fig. 2 Q). Thus, the area occupied by the pericentrin-defined MTOC was less than the tubulin-defined MTOC. Colocalization between the pericentrin-defined MTOC and CIP4, however, was greater in cytolytic conjugates than the colocalization between CIP4 and the tubulin-defined MTOC. This difference was not found in noncytolytic conjugates, as CIP4 colocalization with the MTOC was less (note MTOC enlargement in Fig. S2, available at http://www.jem.org/cgi/content/full/jem.20061893/DC1) and, thus, is consistent with the centrosomal focusing of CIP4 in cytolytic conjugates. As an alternative to these whole-cell area and intensity cell-based colocalization analyses, the colocalization within a constant size region around the MTOC was also performed and defined a similarly increased colocalization between the MTOC and CIP4 in YTS CD2-GFP cells engaged in cytolytic versus noncytolytic conjugates (Fig. S2).

Because CIP4 could have the potential to facilitate interaction between the MTOC and F-actin during formation of
Figure 2. Accumulation of CIP4 with the MTOC at the cytolytic but not noncytolytic IS. The cytolytic IS (A–F) between a YTS CD2-GFP cell and a KT86 cell, and a noncytolytic IS (G–L) between a YTS CD2-GFP cell and a K562 cell, as viewed by DIC microscopy (A and G), as well as confocal microscopy showing fluorescence for α-tubulin (B and H), pericentrin (C and I), CIP4 (D and J), CD2-GFP (E and K), and an overlay (F and L). Although an x-y plane was selected in the z axis to highlight the MTOC, disproportionate accumulation of CIP4 in other x-y planes was not found. (M) Accumulation of CD2, the tubulin-defined MTOC, or CIP4 in >200 conjugates over four experiments at the cytolytic (black bars) and noncytolytic (gray bars) IS. (N) Mean distance of the tubulin- (M to S) or pericentrin-defined (P to S) MTOC from the IS in YTS cells conjugated with KT86 (black bars) or K562 (gray bars) target cells is shown. Distances were greater in noncytolytic conjugates. (O) The area of α-tubulin, the MTOC, and CIP4 was measured in the x-y plane containing the MTOC in YTS cells conjugated with KT86 (black bars) or K562 (gray bars) target cells. The MTOC was defined as the region consisting of accumulated α-tubulin through the exclusion of nonaccumulated α-tubulin using a size and intensity threshold. (P) The percent colocalization calculated among areas is shown for the percentage of α-tubulin that colocalized with CIP4 (T/C), the MTOC that colocalized with CIP4 (M/C), CIP4 that colocalized with the MTOC (C/M), and CIP4 that colocalized with α-tubulin (C/T). Significant differences between the cytolytic (black bars) and noncytolytic (gray bars) synapses are noted. (Q) In YTS cells engaged in cytolytic (black bars) and noncytolytic (gray bars) conjugates, the mean percentage of the pericentrin-defined MTOC that colocalized with the α-tubulin-defined MTOC (P/M), the percentage of the α-tubulin-defined MTOC that colocalized with the pericentrin-defined MTOC (M/P), and the percentage of the pericentrin-defined MTOC that colocalized with CIP4 (P/C) are shown. Error bars represent the SD. *, P < 0.01.
the mature lytic IS, F-actin was evaluated relative to CIP4 and the MTOC after their increased association at the cytolytic IS. NK92 cells were studied in addition to YTS cells to ensure that the findings were not unique to YTS (Fig. 3, A–L). K562 cells are susceptible to NK92 cell–mediated lysis and, thus, were used as physiologic target cells for the NK92 cells. To discern the NK cell from the target cell, YTS cells expressing GFP were conjugated with KT86 cells. Similarly, K562 cells expressing GFP were used to make conjugates with NK92 cells. In each case, conjugates were selected that contained one GFP+ cell paired with one GFP− cell (Fig. 3, B and H). F-actin, the α-tubulin–defined MTOC, and CIP4 accumulated at the majority of synapses in both YTS (Fig. 3, C–F) and NK92 cells (Fig. 3, I–L), though the F-actin accumulation was a more consistent feature (Fig. 3 M). The area occupied by F-actin, the MTOC, and CIP4 was equivalent in the two NK cell types (Fig. 3 N), as was the pattern of colocalization. Specifically, slightly more than half of the total MTOC area contained CIP4 and slightly less than half of the total CIP4 area contained the MTOC (Fig. 3 O). In contrast, approximately one quarter of the total F-actin area contained CIP4 and only one tenth of the total CIP4 area contained

Figure 3. Colocalization of CIP4 with the MTOC and accumulated F-actin at the cytolytic IS. Cytolytic conjugates between a YTS GFP cell and a KT86 cell (A–F), and an NK92 cell and a K562 GFP cell (G–L). DIC images (A and G) and GFP expression in the YTS cell (B), or in the K562 cell (H), distinguish the NK cell from the target cell. Confocal microscopy for F-actin (C and I), α-tubulin (D and J), CIP4 (E and K), and an overlay (F and L) are shown. (M) Molecular accumulations in NK cells from ≥150 conjugates over at least three experiments between NK92 and K562 cells, YTS GFP and KT86 cells, YTS CIP4 and KT86 cells, CIP4 FCH+ and KT86 cells, and CIP4 SH3− and KT86 cells. CIP4 accumulation was not determined for the CIP4-overexpressing cells, as it was diffuse. (N) The total area of F-actin, the tubulin-defined MTOC, and CIP4 measured in ≥15 conjugated NK cells over three experiments between NK92 and K562 cells (black bars), YTS GFP and KT86 cells (light gray bars), and YTS CIP4 and KT86 cells (dark gray bars). CIP4 mutants were not detected with anti-CIP4 mAb. (O) The percent colocalization among the areas in ≥15 conjugated NK cells over three experiments between NK92 and K562 cells (black bars), YTS GFP and KT86 cells (light gray bars), and YTS CIP4 and KT86 cells (dark gray bars) was calculated to demonstrate the percentages of the MTOC colocalized with CIP4 (M/C), of the CIP4 colocalized with the MTOC (C/M), of the F-actin colocalized with CIP4 (A/C), and of the CIP4 colocalized F-actin (C/A). Differences between YTS GFP and NK92 cells were not significant, but those between YTS CIP4 and YTS GFP cells were. Error bars represent the SD. *, P < 0.01.
F-actin. Thus, at the cytolytic IS, CIP4 shared space with a substantial portion of the MTOC and a small amount of F-actin. This raised the possibility that CIP4 could serve a functionally important role in linking the MTOC to a critical region of F-actin at the IS.

**CIP4 or truncated CIP4 overexpression blocks NK cell cytotoxicity by altering cytoskeletal associations and preventing MTOC polarization to the IS**

To investigate the functional importance of CIP4 in cytotoxicity and at the cytolytic IS, we initially took an overexpression approach. We hypothesized that if CIP4 was to serve a role as an important link between the MTOC and F-actin, unregulating its expression would interfere with its physiological role as a linker. Full-length WT CIP4 cDNA or CIP4 cDNA deleted of the nucleotides encoding the first 117 aa encompassing the FCH domain (FCH⁻), or encoding the last 128 aa encompassing the SH3 domain (SH3⁻), were cloned into the pMX-IRES-GFP retroviral vector (Fig. 4 A). These vectors were packaged into ecotropic retroviruses and were used to infect YTS cells stably expressing the ecotropic receptor (22). Transduced cells were identified through GFP expression and sorted by FACS to create cultures stably expressing high or low levels (Fig. 4 B). Anti-CIP4 pAb was used in Western blots of lysates of these cells to confirm the presence of CIP4 protein, and the high GFP-expressing cell lines (YTS-CIP4) had the highest levels of CIP4 protein (Fig. 4 C). The low expressing cells still had levels of CIP4 that were higher than endogenous levels. The pAb failed to recognize the FCH⁻, and the mAb failed to recognize either mutant construct (unpublished data). Overexpressed WT CIP4 could be detected using mAbs by intracellular FACS.
and was present at substantially higher levels than YTS, NK92, or ex vivo NK cells (Fig. 1, C and D).

Because NK cell cytolytic activity requires the coordinated and sequential function of both actin and microtubules, we determined the impact of CIP4 overexpression by initially measuring cytotoxicity. The lytic activity of all six new cell lines was measured against KT86 target cells and was compared with YTS cells expressing levels of GFP similar to those found in the high CIP4-expressing lines. In contrast to cells expressing GFP alone, cytotoxic function was inhibited by overexpression of the WT or mutant CIP4 constructs (Fig. 4 D). This was not caused by decreased CD28 expression (unpublished data). Inhibition of cytotoxicity by CIP4 overexpression was not absolute but was proportional to the level of transgene expression and did not require the simultaneous presence of the FCH or SH3 domains. This result suggested that unregulated expression of CIP4 results in an abnormal role for this protein that interferes with the normal physiological process of cytotoxicity.

To determine how CIP4 overexpression interferes with cytotoxicity, CIP4-overexpressing cells conjugated with KT86 target cells were evaluated by confocal microscopy to study cytoskeletal reorganization and CIP4 localization. YTS cells overexpressing CIP4 were identified in conjugates with KT86 cells through expression of GFP via the IRES in the transduced construct (Fig. 4 F). In cells overexpressing WT CIP4, the tubulin-defined MTOC generally failed to polarize to the IS, but F-actin still accumulated (Fig. 4, G and H). Interestingly, in WT CIP4-overexpressing cells, CIP4 was no longer primarily found with the MTOC but was now found in the cell cortex, as seen by colocalization with F-actin. The decreased polarization of the MTOC to the IS was a consistent feature of CIP4 overexpression even in the absence of the FCH or SH3 domains, as all overexpressing cells had less than one third the polarization of the tubulin-defined MTOC compared with cells expressing GFP alone (Fig. 3 M). Similar to the WT CIP4-expressing cells, each of the mutant-expressing lines maintained the ability to accumulate F-actin at the IS (Fig. 3 M and Fig. S3, available at http://www.jem.org/cgi/content/full/jem.20061893/DC1), suggesting that the impact of CIP4 overexpression was downstream of F-actin reorganization. As perforin and lytic granules polarize to the activating NK cell IS with the MTOC (12, 13), perforin was evaluated in YTS cells overexpressing each of the CIP4 constructs and did not appear disrupted in its normal proximity to the MTOC (Fig. S3). Thus, the movement of perforin-containing lytic granules relative to the MTOC would not be expected to be impaired by CIP4 overexpression. Instead, these data indicate a more specific effect of CIP4 on MTOC localization.

Because actin accumulation at the IS appeared intact, a potential mechanism by which CIP4 overexpression could interrupt MTOC polarization and cytotoxicity might be by occupying physiological binding sites in the F-actin network. Quantitative analysis of confocal micrographs demonstrated that although the area occupied by CIP4 after overexpression increased twofold (Fig. 3 N), the colocalization between F-actin and CIP4 increased threefold (Fig. 3 O), and a majority of F-actin colocalized with CIP4. There was a slight increase in the area of the tubulin-defined MTOC that colocalized with CIP4, but the relative amount of the CIP4 that colocalized with the MTOC was decreased by 49%. Similarly, the relative amount of CIP4 colocalized with F-actin increased after overexpression, and the CIP4 was almost equally
divided in colocalization with the MTOC and F-actin. Thus, after overexpression, CIP4 appears to occupy nonphysiologic space in the F-actin cortex, potentially interfering with the normal physiologic role of the CIP4 that is associated with the MTOC.

To confirm this association biochemically, CIP4 was immunoprecipitated from WT CIP4-overexpressing YTS, parental YTS, and ex vivo NK cells. In lysates of the overexpressing cells, CIP4 coimmunoprecipitated with both α-tubulin and WASp (as predicted by the microscopy data), whereas in parental YTS and ex vivo NK cells, CIP4 preferentially coimmunoprecipitated with α-tubulin (Fig. 5 A). This demonstrates that overexpressed CIP4 was nonphysiologic in its association with WASp but that it had the capacity to link elements that function in the F-actin cortex with α-tubulin. To understand if the association between CIP4 and α-tubulin in YTS cells could be indicative of an association with microtubules, YTS lysates were cleared of debris and exposed to stabilized microtubules. Microtubules removed CIP4 from YTS lysates and led to CIP4 concentration in the microtubule precipitate (Fig. 5 B). Collectively, these data show that CIP4 is a microtubule-associated protein in YTS cells but is largely not associated with WASp under resting conditions.

**Figure 6.** Localization of CIP4 in normal and WASp-deficient ex vivo NK cells. An NK cell (top) and target cell (bottom) in conjugates between a normal donor ex vivo NK cell and a K562 target cell (A–E), as well as a WAS-patient ex vivo NK cell and a K562 target cell (F–J), using DIC microscopy (A and F). F-actin (B and G), α-tubulin (C and H), CIP4 (D and I), and an overlay (E and J) were detected using confocal microscopy. (K) Molecular accumulation at the IS in >150 NK cells conjugated with K562 target cells in at least three experiments are shown using normal donor (black bars) or WAS-patient (gray bars) NK cells. (L) The area of F-actin, the α-tubulin–defined MTOC, and CIP4 was measured in >15 conjugated NK cells over three experiments between K562 target cells and normal donor (black bars) or WAS-patient NK cells (gray bars). (M) The percentage of the MTOC that colocalized with CIP4 (M/C), CIP4 that colocalized with the MTOC (C/M), F-actin that colocalized with CIP4 (A/C), and CIP4 that colocalized with F-actin (C/A) was calculated for the measured areas in the conjugated normal donor and WAS-patient NK cells. Decreases in WAS-patient NK cells were significant. *, P < 0.01.

**WASp is required for the activation-induced physiologic associations of CIP4**

To extend these results to ex vivo NK cells and to evaluate the role for WASp in greater detail, CIP4 was studied in NK cells prepared from healthy donors or a patient with WAS who had no WASp protein. Like YTS and NK92 cells, in conjugates formed between ex vivo NK cells and susceptible target cells, CIP4 and the tubulin-defined MTOC polarized to the IS near the region of accumulated F-actin (Fig. 6, A–E). A similar result was obtained in ex vivo NK cells using a pericentrin-defined MTOC (Fig. S4, available at http://www.jem.org/cgi/content/full/jem.20061893/DC1). In contrast, when ex vivo NK cells were deficient in WASp, there was reduced F-actin accumulation, as well as reduced polarization of the MTOC and CIP4 (Fig. 6, F–K). In the WAS patient NK cells, the CIP4 appeared to be more diffusely located throughout the cytoplasm and was typically not seen in the cell cortex. Compared with control cells, the total area occupied by F-actin and the MTOC was reduced in WAS-patient NK cells (Fig. 6 L). The area occupied by CIP4, however, was equivalent to that found in control cells. As expected, a lower percentage of the CIP4 area in WAS-patient cells colocalized with the reduced areas of either F-actin or the MTOC (Fig. 6 M). Surprisingly, despite the
similar area of CIP4 in WAS-patient cells, the percentage of the MTOC or F-actin that colocalized with CIP4 was less than that found in healthy donor ex vivo NK cells. Thus, WASp was required to enable the normal physiologic localization and associations of CIP4.

The association between CIP4 and WASp is specific to NK cell activation

Because WASp was required for the physiologic associations of CIP4 but CIP4 only coimmunoprecipitated with WASp after CIP4 overexpression (Fig. 5A), we hypothesized that NK cell activation enables CIP4 function and, specifically, the association between WASp and CIP4. To evaluate this, we needed to activate NK cells in the absence of target cells, because the target cell can serve as a source of both WASp and CIP4. This would make it impossible to discern whether the molecule being studied was contributed by the effector or the target cell. Furthermore, the presence of target cell protein in a cell lysate may be of a distinct activation state and could thus dilute or affect results. To activate YTS and ex vivo NK cells, activation receptors were clustered by placing cells on antibody-coated surfaces. Anti-CD28 was used for YTS cells and anti-NKp30 was used for ex vivo NK cells, as these antibodies have been shown to trigger cytotoxicity and activation signaling (21, 23). Cells were incubated for 30 min on glass slides (for microscopy) or plastic wells (for biochemical analysis) coated with the antibody. Surfaces coated with either IgG or media were both used as controls. To confirm that the activating signal had an effect on F-actin, the MTOC, and CIP4, cells were evaluated using laser scanning confocal microscopy and scanning through the x-z axis to visualize their contact with the antibody. Both YTS cells and ex vivo NK cells demonstrated a preferential accumulation of F-actin at, as well as a polarization of the tubulin-defined MTOC and CIP4 to, the activating antibody–coated surface when compared with the IgG-coated surface (Fig. 7, A–D). The IgG-coated slide, however, did induce some F-actin accumulation in ex vivo NK cells at the glass surface when compared with slides not coated with antibody (unpublished data). This was presumably caused by the Fc receptor present on ex vivo NK cells, which was absent on YTS cells. This is consistent with data demonstrating IS formation without MTOC polarization after CD16 ligation in NK cells (24). Despite this, cells with F-actin, the MTOC, and CIP4 accumulated at the activating mAb–coated glass surface more than doubled in number when compared to the IgG-coated glass surface (n ≥ 100 adhered cells over three experiments). The colocalization between the CIP4 and the MTOC also

Figure 7. Alteration in association of endogenous CIP4 after NK cell activation. YTS cells were incubated on slides coated with IgG (A) or anti-CD28 (B) and ex vivo NK cells on slides coated with IgG (C) or anti-NKp30 (D) for 30 min and evaluated via confocal microscopy in the x-z plane. The arrowheads show the plane of the slide. Fluorescence demonstrating F-actin (cyan), α-tubulin (blue), and CIP4 (red) is shown. (E) 10^6 YTS cells were incubated in wells containing IgG, anti-CD28 mAb, or media for 30 min and then lysed. CIP4 was immunoprecipitated using mAb anti-CIP4 and probed for WASp and CIP4 by Western blotting. Immunoprecipitation with nonspecific mIgG was performed as a control. (F) 10^7 ex vivo NK cells were lysed after a 30-min incubation in wells coated with IgG or anti-NKp30 mAb, and CIP4 was immunoprecipitated using mAb anti-CIP4. WASp and CIP4 were detected in immunoprecipitates by Western blotting. Immunoprecipitation with nonspecific mIgG was performed in parallel as a control. Blots represent at least three independent results.
increased after YTS cells were exposed to anti-CD28 or ex vivo NK cells were exposed to anti-NKp30. Specifically, when activated cells were quantitatively analyzed and compared with values obtained from control-treated cells, the increase in colocalization between CIP4 and the MTOC was 27% in YTS cells and 24% in ex vivo NK cells (≥15 cells over three separate experiments quantitatively evaluated for each condition). The increased colocalization between CIP4 and the MTOC after activating receptor signaling suggested that the CIP4 aggregates in the MTOC after activation. The colocalization between CIP4 and F-actin was also increased in cells on the activating antibody–coated surface (15% in YTS cells and 21% in ex vivo NK cells). This suggested that CIP4 aggregated with the MTOC associates with the F-actin cortex after NK cell activation.

To biochemically discern an activation-induced association, CIP4 was immunoprecipitated from YTS cells that were exposed to plastic coated with IgG, media, or anti-CD28. The presence of WASp was evaluated in the immunoprecipitate by Western blotting (Fig. 7 E). Similar experiments were performed using ex vivo NK cells incubated on plastic surfaces coated with IgG or anti-NKp30 (Fig. 7 F). In either case, although CIP4 levels did not increase, there was an increased association between WASp and CIP4 after activation. Thus, after NK cell activation, CIP4 has an increased association with WASp and, therefore, has the potential to link actin branches in the cell cortex to the MTOC to serve a role in localizing the MTOC to the IS.

CIP4 expression is required for NK cell cytotoxicity and MTOC polarization

To demonstrate a role for endogenous CIP4 in MTOC polarization and cytotoxicity, small interfering RNA (siRNA) specific for CIP4 was used. siRNA was introduced into YTS cells via nucleofection, and siRNA directed against GAPDH- or mock-transfected cells were used as controls. There were no differences found between GAPDH siRNA–transfected and mock-transfected cells for any of the parameters evaluated except for GAPDH expression (unpublished data). Introduction of siRNA specific for CIP4 reduced expression of CIP4 by 47.4 ± 23.8% (P < 0.001) in YTS cells over nine independent experiments, as determined by Western blotting (Fig. 8 A). This reduction was specific to CIP4 siRNA, as GAPDH–specific siRNA did not reduce CIP4 expression. Similar results were obtained using a FACS analysis for CIP4 in siRNA nucleofected cells (unpublished data). Cytolytic activity was reduced by a mean of 42% across different effector/target cell ratios after CIP4 siRNA nucleofection when compared with GAPDH siRNA (Fig. 8 B). To determine if this was caused by an effect of reduced CIP4 expression on MTOC polarization to the IS, YTS cells nucleofected with either CIP4 or GAPDH siRNA were conjugated with KT86 target cells and evaluated via confocal microscopy. Although the accumulation of F-actin at the IS was comparable in both, there was a near 50% reduction in polarization of the tubulin–defined MTOC to the IS in cells nucleofected with CIP4 siRNA. This demonstrated that IS formation had occurred in the presence of reduced CIP4 levels but that MTOC polarization was impaired. Therefore, the role of CIP4 in NK cell function is most likely in facilitating the localization of the MTOC to the IS, which is required for polarization of the lytic granules and cytotoxicity.

Relation of CIP4 function to Cdc42 activation

Because the activity of Cdc42 is required for the polarization of the MTOC to the IS (14), we wanted to determine the relation of CIP4 to Cdc42 activity. We assumed that CIP4 would function downstream of Cdc42, as interruption of Cdc42 activity interrupts both F-actin accumulation and MTOC polarization at the IS, whereas interference with CIP4 function only affected the latter. Two approaches were taken to determine if the requirement for CIP4 function followed...
Collectively, these results demonstrated that CIP4 siRNA affects MTOC polarization and NK cell function even in the presence of active Cdc42.

As a second approach, active Cdc42 was isolated from anti-CD28-activated YTS cells (Fig. 9 O). In this case, active Cdc42 was present in activated YTS cells nucleofected with GFP, but there was more in YTS cells nucleofected with Cdc42V12, demonstrating the specificity of the assay. To evaluate the activation of Cdc42 in the presence of reduced CIP4, YTS cells were nucleofected with only CIP4 siRNA or GAPDH siRNA, activated, lysed, and active Cdc42 isolated. CIP4 siRNA did not reduce the amount of active Cdc42, suggesting that Cdc42 activation could occur independently of normal CIP4 expression. As a final consideration, YTS cells were compared with YTS cells overexpressing WT CIP4, which were inhibited in MTOC polarization and cytotoxicity. Again, the amount of active Cdc42 that could

Figure 9. CIP4 function in MTOC polarization relative to Cdc42 activation. The cytolytic IS between a KT86 cell and a YTS cell nucleofected with constitutively active Cdc42V12-GFP and GAPDH siRNA (A–F) or CIP4 siRNA (G–L) are shown using DIC microscopy (A and G), as well as confocal microscopy showing fluorescence for α-tubulin (B and H), pericentrin (C and I), CIP4 (D and J), Cdc42V12-GFP (E and K), and an overlay (F and L). (M) The mean percentage of the pericentrin-defined MTOC that colocalized with the tubulin-defined MTOC (P/M) and the percentage of the α-tubulin-defined MTOC that colocalized with the pericentrin-defined MTOC (M/P) in YTS cells nucleofected with Cdc42V12-GFP and GAPDH siRNA (black bars) or Cdc42V12-GFP and CIP4 siRNA (gray bars), and conjugated with KT86 target cells, are shown. (N) Mean distance in micrometers of the pericentrin-defined MTOC to the IS in YTS cells nucleofected with Cdc42V12-GFP and GAPDH siRNA, or Cdc42V12-GFP and CIP4 siRNA. The increase in cells receiving CIP4 siRNA was significant, * P < 0.01. (O) Active Cdc42 pull down and Cdc42 Western blot from YTS cells activated with immobilized anti-CD28. Before activation, cells were nucleofected with GFP or Cdc42V12-GFP (top), or nucleofected with GAPDH siRNA or CIP4 siRNA (middle). Activated YTS cells were also compared with activated YTS cells overexpressing WT CIP4 (bottom). Error bars represent the SD.

or preceded Cdc42 activation. The first was to nucleofect YTS cells with a previously described constitutively active Cdc42V12 construct (15, 25) and GAPDH or CIP4 siRNA. When these cells were conjugated with KT86 target cells to form a cytolytic IS, MTOC localization to the IS occurred normally in those that had received GAPDH, but not CIP4, siRNA (Fig. 9, A–L). The colocalization between the pericentrin and α-tubulin–defined MTOC was not altered by CIP4 siRNA, suggesting that the integrity of the MTOC was not affected (Fig. 9 M). However, the polarization of the MTOC as measured by the mean distance of the pericentrin–defined MTOC to the IS was decreased after CIP4 siRNA nucleofection in the presence of constitutively active Cdc42V12 (Fig. 9 N). Concomitantly, the cytolytic activity of Cdc42V12 nucleofected cells that had received CIP4 siRNA compared with those that had received Cdc42V12 and GAPDH siRNA was reduced by 34 ± 9% across effector/target cell ratios.
be isolated from activated YTS cells was not reduced by CIP4 overexpression. Thus, Cdc42 activation occurred in cells with interrupted CIP4 function, suggesting that CIP4 functions downstream of Cdc42 to induce MTOC polarization to the IS and cytotoxicity.

**DISCUSSION**

The cytolytic IS forms and functions in a series of stages. Some of these stages are linear in sequence and, thus, define key regulatory steps on which function of the IS depends. Previously, we found that F-actin and cell-surface receptor reorganization at the SMAC do not require microtubule function and precede the delivery of cytolytic granules to the IS (10). Lytic granule traffic, however, required both microtubule integrity and WASp-dependent F-actin function. Thus, these two events are sequential, suggesting the utility of potential signaling and structural linkages between them. Exact delivery of lytic granules is most likely a critical component of directed cytotoxicity and is consistent with work defining a specific cytolytic secretory domain at the edge of the central SMAC within the IS (26). This delivery of lytic granules to the IS has more recently been shown to be directed by the MTOC itself (11). Thus, localization of the MTOC is critical and may benefit from a specific connection between F-actin at the SMAC and microtubules. Because WASp is localized with F-actin at the pSMAC, including at the border of the secretory domain (16), a link between microtubules and WASp could serve a critical function in directing the MTOC and associated lytic granules to the secretory domain of the cytolytic IS. For these reasons, we investigated CIP4 for its potential to link WASp and tubulin as a critical regulator of the cytolytic immunological IS.

CIP4 was discovered because of its ability to bind to Cdc42 (17) and WASp (18) through two separate efforts using the yeast two-hybrid system. Overexpression of WASp and CIP4 in fibroblasts led to association of WASp with microtubules mediated by the FCH domain of CIP4 (18). It was subsequently appreciated that CIP4 deficient of its FCH or SH3 domains microinjected into macrophages could interfere with the formation of microtubule and WASp-dependent podosome formation (19). Collectively, these studies suggested the potential for CIP4 to provide relevant associations with both WASp and microtubules.

To determine if the CIP4 system was relevant to NK cell biology and potentially directed secretion, we initially identified that CIP4 was expressed in NK cells. CIP4 was diffusely present in cells that were not activated or in noncytolytic conjugates, as determined using confocal microscopy. After cell activation by either a target cell or NK cell activation receptor signaling, however, CIP4 accumulated with the MTOC (as defined either by α-tubulin or pericentrin) at the IS (Figs. 2, 3, 6, 7, and 9; and Figs. S1, S2, and S4). This suggested that CIP4 functioned by joining the MTOC after cell activation. CIP4, however, could be coimmunoprecipitated with α-tubulin even from resting NK cells in the absence of target cells (Fig. 5 A). This raised the possibility of a model in which CIP4 in resting cells is complexed with free tubulin, or with tubulin in individual microtubules, but is then focused to the MTOC during cell activation. Support for this model was consistent with CIP4 having reduced colocalization with the MTOC in resting NK cells or NK cells in non-cytolytic conjugates, but was suggested by some α-tubulin still being colocalized with CIP4. Moreover, the resolution of confocal microscopy is likely to underestimate the actual association between individual microtubules, or free tubulin, and CIP4. Also, as we have only evaluated α-tubulin, there may be substantial associations between other forms of tubulin and CIP4, resulting in further underestimation.

Focusing CIP4 to the MTOC may have at least two functions. One would be in the actual polarization of the MTOC to the IS and the other in MTOC anchoring at the IS. The first of these two would require that CIP4 join the MTOC after cell activation but before MTOC polarization. A requirement for CIP4 in polarization of the MTOC is highlighted by the siRNA results and is further suggested by the overexpression studies in which polarization of the MTOC to the IS was decreased. Although MTOC polarization was reduced in cells overexpressing WT CIP4, this same result was also found in cells overexpressing CIP4 deleted of the FCH domain (Fig. 3 M). As this molecule is lacking its tubulin-interacting region, an alternative purpose to focusing CIP4 to the MTOC is suggested. Specifically, CIP4 may serve to anchor the MTOC at the appropriate location at the SMAC to allow directed delivery of the lytic granules. This anchoring function could be facilitated through the interaction of MTOC-associated CIP4 with WASp present in the F-actin cortex. Support for an anchoring function is provided by WASp-deficient NK cells, in which localization of the MTOC to the IS did not occur (Fig. 6). Alternatively, this result could be a feature of defective NK cell activation in the absence of WASp. In particular, ex vivo WAS NK cells demonstrate reduced adhesion to target cells (27), and those that have conjugated fail to cluster CD11a or CD11b at the IS (10). Thus, WASp-deficient NK cells may be receiving a weaker activation signal for MTOC polarization because of reduced integrin signaling. Our CIP4 overexpression studies, however, are also consistent with an MTOC-anchoring role for CIP4. Overexpressing CIP4 or CIP4 lacking its FCH domain would be capable of interacting with WASp without being associated with tubulin and, thus, could interfere with the interaction of tubulin-associated CIP4 with WASp. In this light, immunoprecipitation of CIP4 in cells overexpressing the WT construct yielded only slightly more tubulin than did CIP4 immunoprecipitated from parental cells (Fig. 5). Conversely, overexpression of the SH3-deleted construct of CIP4 may allow association with tubulin in the MTOC but prevent an anchoring function to WASp in the SMAC. Results obtained using CIP4 siRNA (Fig. 8) do not differentiate between these two models and only demonstrate a requirement for CIP4 in MTOC localization to the IS. As the MTOC is very dynamic in cytotoxic lymphocytes (28, 29), the ability to provide a strong anchor at the SMAC may be useful in
maintaining the polarity of the MTOC. This, in turn is likely to be valuable in directing large numbers of lytic granules through the IS onto the target cell.

Given that CIP4 is infrequently found in the F-actin cell cortex in resting NK cells or in NK cells in noncytolytic conjugates, a specific mechanism for facilitating an interaction between CIP4 and WASp must be in place. Regulation could be mediated through the modification of CIP4, WASp, or both. Because CIP4 was associated with tubulin in resting cells (Fig. 5), the interaction with WASp may represent the critical step. Regulation of WASp has been extensively studied (for review see reference 30). WASp exists in an autoinhibited conformation in which its verprolin, cofilin, acidic domain and its GTPase binding domains are in contact, obscuring key elements required for F-actin branching (31). WASp autoinhibition is released after activation by the phosphorylation of key tyrosines. In unstimulated NK cells, where a majority of WASp would be expected to be autoinhibited, we found minimal WASp associated with CIP4 after CIP4 immunoprecipitation. As NK cell activation increased the association between CIP4 and WASp (Fig. 7, E and F), we propose that CIP4 preferentially interacts with activated WASp. The ability to detect WASp in the immunoprecipitates from cells overexpressing CIP4 could reflect the ability of a greatly increased number of CIP4 molecules to interact with the relatively small amount of activated WASp that exists in resting cells. Alternatively, CIP4 overexpression could influence the activation state of WASp specifically. An interaction between CIP4 and activated WASp, however, would present a useful means for specifically anchoring the MTOC at the SMAC, where WASp is functioning to rearrange F-actin. Similarly, it would present a useful means of releasing the MTOC once signaling is terminated and WASp reverts to its autoinhibited state. Although further study of MTOC dynamics will be required to specifically prove this hypothesis, in preliminary experiments we did find phosphorylated WASp associated with CIP4 after physiologic NK cell activation (unpublished data). As WASp is expressed only in hematopoietic cells, this would represent a hematopoietic cell–specific mechanism for positioning and/or anchoring the MTOC. CIP4 does interact with the more ubiquitously expressed WASp homologue neural WASp (32, 33) and, thus, can serve a more general role in MTOC positioning. However, because neural WASp does not completely compensate for WASp deficiency in NK cells (16, 27), the interaction between WASp and CIP4 may have a specific role in controlling MTOC positioning in the immune system.

Not all lytic granules, however, require the MTOC and microtubule function for activation-induced polarization to the IS. Some lytic granules are “peldocked” with the NK cell membrane (34) and are most likely poised for rapid release onto target cells without the need for substantial cytoskeletal rearrangement. This may in part explain why the CIP4-overexpressing cells, or cells nucleofected with CIP4 siRNA, still mediate some cytotoxicity even after having reduced MTOC polarization (Fig. 4 D and Fig. 8 B). Alternatively, cytotoxicity in these cells was mediated only by the subset in which MTOC polarization occurred. It is more likely that CIP4 as well as MTOC polarization and anchoring are likely to only be required for a portion of NK cell cytotoxicity. Specifically, they would be required for that which depends on granule delivery from elsewhere in the cell to the IS. For this reason, we also considered it important to use ex vivo NK cells that had not been cultured or otherwise prestimulated in the experiments. Activation signals, such as those commonly provided when culturing NK cells, could increase the percentage of lytic granules that are predocked at the cell membrane and reduce reliance on the microtubule-directed transport of granules. Alternatively, NK cell activation signals may induce different mechanisms of enabling MTOC anchoring at the IS. In this light, NK cell activation with IL-2 did not increase the cellular levels of CIP4 but did increase MTOC polarization (unpublished data). Even without activation, however, any linkage between the MTOC and F-actin in the SMAC may be redundant. As evidence, the residual MTOC polarization and cytotoxicity in the CIP4-overexpressing and knockdown cells may reflect alternative mechanisms for MTOC anchoring at the IS. A candidate alternate protein is LIS1, which facilitates the attachment of microtubules to the cell cortex in Dicyostelium through its ability to associate with dynemin and Rac1 (35). In contrast to CIP4, however, LIS1 is inherently associated with the centrosome and is also involved in actin reorganization. Our earlier experiments suggest the existence of a linking protein that is not required for actin dynamics, because microtubule integrity (as disrupted by colchicine treatment) is not required for actin accumulation at the SMAC (10).

Finally, as CIP4 can also interact with Cdc42 (17) and Cdc42 is required for MTOC reorientation to the IS (14), the function of CIP4 could have been through facilitating Cdc42 activity. Because interfering with Cdc42 activity reduces F-actin reorganization at the IS (14) and lytic granule polarization to the IS occurs after F-actin has accumulated (10), we focused on a protein that would block MTOC polarization without interfering with F-actin accumulation at the SMAC. CIP4 functions downstream of Cdc42, as the effect of reducing CIP4 in blocking MTOC reorientation was still apparent in the presence of constitutively active Cdc42 (Fig. 9). Furthermore, after NK cell activation, active Cdc42 levels were unaffected by CIP4 siRNA or CIP4 overexpression (Fig. 9). Thus, it is likely that Cdc42 function represents another early requirement in facilitating the signals leading to MTOC reorientation more akin to WASp, which was required for both F-actin accumulation and MTOC polarization (Fig. 6). It is possible, however, that CIP4 in the MTOC may use Cdc42 as an additional anchor to facilitate MTOC positioning.

As directing lytic granules to the IS is a required component of the specific lysis of target cells, the ability to direct and anchor the MTOC at the IS can be an essential step in effective lytic function. Polarizing and anchoring the MTOC
can allow for the delivery of substantial numbers of lytic granules to the IS and enable maximal exocytosis of lytic granules. CIP4 is a protein that participates in this process first through being focused to the MTOC and then by interacting with WASp. CIP4 has the functional capacity to link actin and microtubule networks in NK cells downstream of Cdc42, and its appropriate expression is required for polarization of the MTOC to the cytolytic IS. CIP4 presents an important paradigm in NK cells that may apply to other immune cells, by which immunological activity can be enabled through functional molecular linkage between the SMAC and MTOC.

MATERIALS AND METHODS

NK cell preparation, NK cell lines, and cellular evaluation. Leukocyte-enriched blood was obtained from volunteer donors and was used to prepare ex vivo NK cells by negative depletion using the human NK cell isolation kit II (Miltenyi Biotec). Ex vivo NK cells from a patient with a G252A point mutation in the acid core facility of the Children’s Hospital of Philadelphia. Other overhangs were provided by D.M. Stewart and D.L. Nelson (National Institutes of Health, Bethesda, MD). Steal and Notl restriction sites were added to these constructs via overhanging PCR amplification to allow cloning into pTOPO zero blunt (Invitrogen) and, subsequently, the bicistronic retroviral expression vector pCMV-cis/HS (38). All human samples were obtained after informed patient or parental consent and were used with approval of the institutional internal review board for the protection of human subjects of the Children’s Hospital of Philadelphia or the Children’s Hospital Boston. The immortalized NK cell lines YTS, YTS expressing the mouse ecotropic receptor (YTSeco), YTS GFP expressing GFP, YTS CD2-GFP expressing a CD2-GFP fusion protein, and NK92, as well as the MHC class I-negative K562 erythroblasts target cell line, have all been previously used and described (10, 36). GFP-expressing K562 cells (K562-GFP) were generated by transfection of K562 cells with a GFP expression vector (CLONTECH Laboratories, Inc.) containing a neomycin resistance gene and GFP maintained via neomycin selection. K562 cells stably expressing CD86 (KT86 cells) were created with CD86 cloned from cDNA (prepared from matured dendritic cells), which was then cloned into lentivector vectors, as previously described (37). Lentivirus-transduced K562 cells were sorted for high GFP expression, and the KT86 line was derived from a single-cell clone. NK cell cytotoxicity against KT86 target cells was evaluated by a 4-h 

Immunoprecipitation and Western blotting. Cells were lysed in ice-cold lysis buffer (300 mM Tris-Cl [pH 7.5], 100 mM NaCl, 15 mM EDTA, 0.1% Triton X-100 and protease inhibitor cocktail [Roche]). For Western blotting of total lysate, 10 μg of protein per sample was separated on a 4–12% Bis-Tris density gradient gel (Invitrogen) and transferred to polyvinylidene fluoride membranes, which after blocking with 3% BSA and 140 mM NaCl in Tris-buffered saline (TBS) were incubated with mouse anti-CIP4 mAb clone 21 (BD Biosciences). Bound antibody was detected with peroxidase-conjugated light chain–specific goat anti–mouse IgG (Jackson ImmunoResearch Laboratories) and an enhanced chemiluminescence detection system (GE Healthcare). Arrowheads shown in Western blot images denote the location of molecular mass markers. Membranes were stripped in 0.2 M glycine (pH 2.5), 0.05% Tween 20, and 140 mM NaCl in TBS at 50°C for 30 min, blocked with 3% BSA, and reprobed with rabbit anti-WASP pAb (U.S. Biologicals) or rabbit antitubulin pAb (U.S. Biologicals), followed by peroxidase-conjugated light chain–specific goat anti–rabbit IgG (Jackson ImmunoResearch Laboratories). In gene silencing experiments, GAPDH was detected using mouse anti-GAPDH clone ID4 (Imgenex). For immunoprecipitation, lysates were normalized for either protein content or cell number and precleared with 10 μg/ml of mouse mAb clone MOPC21 (BD Biosciences) and protein G agarose beads (Invitrogen). The supernatant was incubated with 20 μg/ml anti-CIP4 mAb or nonspecific mAb IgG MOPC21, followed by protein G agarose beads, and the immunoprecipitate was separated on density gradient gels, followed by Western blotting. For preactivation of cells through CD28 or NKp30, untreated polystyrene sixwell plates (Nunc) were coated with 5 μg/ml of mAb anti-CD28 clone 9.3 (a gift of J. Riley, University of Pennsylvania School of Medicine, Philadelphia, PA) or anti-NKp30 clone 22S (Beckman Coulter), and were washed and incubated with 10 μg YTS or 105 ex vivo NK cells for 30 min at 37°C. Wells coated with mAb MOPC21 were used as a control.

Microtubule binding protein spin-down assay. The assay was performed using a commercially available reagent set according to the manufacturer’s recommendations (Cytoskeleton, Inc.). In brief, microtubules were generated from purified tubulin and stabilized with 200 μM Taxol. YTS cells were lysed in 20 mM ice-cold Heps (pH 7), 2 mM EGTA, and 0.5% Triton X-100 and with protease inhibitor cocktail for 1.5 h, and were cleared of debris by centrifugation at 9,300 g. The cleared lysate was then incubated for 30 min at room temperature in the presence or absence of 10 nM of stabilized microtubules. Microtubules and associated proteins were precipitated by centrifugation at 100,000 g of protein per sample was separated on a 4–12% Tris-buffered saline (TBS) were incubated with mouse anti-CIP4 mAb or nonspecific mAb IgG MOPC21, followed by protein G agarose beads, and the immunoprecipitate was separated on density gradient gels, followed by Western blotting. For preactivation of cells through CD28 or NKp30, untreated polystyrene sixwell plates (Nunc) were coated with 5 μg/ml of mAb anti-CD28 clone 9.3 (a gift of J. Riley, University of Pennsylvania School of Medicine, Philadelphia, PA) or anti-NKp30 clone 22S (Beckman Coulter), and were washed and incubated with 10 μg YTS or 105 ex vivo NK cells for 30 min at 37°C. Wells coated with mAb MOPC21 were used as a control.

Confocal microscopy and image analysis. Conjugates between NK cells and target cells at a 2:1 ratio were formed in suspension for 15 min and adhered to poly-l-lysine–coated glass slides (PolyPrep; Sigma-Aldrich) for 15 min, all at 37°C, as previously described (10). In experiments evaluating NK cell activation without target cells, poly-l-lysine–coated glass slides were coated with IgG MOPC21 as a control, or anti-CD28, or anti-NKp30 by overnight incubation with 5 μg/ml of antibody in PBS. Slides were washed, and NK cells were incubated on the slide in media for 30 min at 37°C, after which nonadherent cells were rinsed. In either case, cells adhering to the slide were fixed and permeabilized with 4% formaldehyde, 0.1% saponin, and 0.1% Triton X-100 in PBS for 15 min, rinsed, and incubated for 1 h with anti-CIP4 mAb or nonspecific IgG clone MOPC21 (as a control). Slides were rinsed, incubated for 1 h with Alexa Fluor 647–conjugated highly cross-adsorbed goat anti-mouse (Invitrogen), rinsed again, and incubated with 10% heat-inactivated mouse serum (Sigma-Aldrich) for 30 min to block nonspecific binding. Slides were rinsed and incubated with biotinylated anti–α-tubulin mAb (Invitrogen) or biotinylated mouse IgG control (BD Biosciences) for 1 h, followed by additional rinsing and incubation with streptavidin–conjugated Pacific blue 405 (Invitrogen) and Alexa Fluor 568–conjugated phallolidin (Invitrogen). Slides were rinsed and covered with
0.15-mm glass coverslips (VWR Scientific) using Prolong Antifade reagent (Invitrogen) or Vectashield with DAPI. Other antibodies used for microscopy, where specified in the figures, include pAb rabbit anti-α-tubulin (Abcam), pAb antipericentrin (Abcam), Alexa Fluor 568–conjugated anti-rabbit IgG (Invitrogen), FITC anti-rabbit IgG (Jackson ImmunoResearch Laboratories), FITC anti-mouse IgG (Jackson ImmunoResearch Laboratories), and mAb FITC antiperform (Becton Dickenson). All antibodies were used in the range of 1–20 μg/ml and were diluted in saponin-containing buffer. For each experiment, cells stained with individual fluorophores and controls for individual fluorophores were included as controls to allow for proper spectral separation and/or adjustment of the microscope.

Cell conjugates were visualized by using either a laser scanning confocal microscope (DMI RE2; Leica) or a spinning-disc confocal microscope (IX81 DSU; Olympus) via scanning through the x-y plane. Detection settings were adjusted so that a control-stained sample was uniformly negative and experimentally stained samples were not saturating or bleeding into other channels. The effector and target cells in the conjugate were confirmed either by size (for eX vivo NK cells) or by the presence or absence of GFP expression. The area of accumulation of fluorescent molecules in the NK cell was examined using the threshold of ≥40% of unaccumulated fluorescence intensity for evaluations of CD2 or F-actin (a number based on previously applied assumptions) (10, 28). Accumulation of CD2 or F-actin at the IS consisted of an increased area of accumulated CD2 or F-actin at the IS relative to other cell-surface or cortical regions. Accumulated areas of CIP4, α-tubulin, and pericentrin were identified using the same threshold approach but were often not found at the IS or in the cell cortex. To determine the area occupied by an accumulated fluorescence intensity and colocalization between areas of different fluorescence, images were analyzed using the Volocity software package classification module (Improvision). The area occupied by each accumulated fluorophore only in the effector cell was measured. Areas containing two accumulated fluorophores were also measured and compared with the area occupied by the single fluorophore to calculate the percentage colocalization. The MTOC was defined either as the accumulated area of α-tubulin using a pixel-size threshold, to eliminate measurement of isolated microtubules, or as the accumulated area of the centrosomal protein pericentrin. As an alternative approach to evaluating colocalization with the MTOC, fluorescence intensities of the individual pixels that were over background levels in a 5-μm² region enclosing the MTOC were measured in some experiments, as noted in the figures. In experiments evaluating NK cell activation by a coated glass surface, scanning was performed using the Leica microscope scanning in the x-z plane to completely visualize the interface with the glass, and analysis was performed as for the conjugates. Polarization of the MTOC to the IS was defined as either there being at least some degree of colocalization of the α-tubulin–defined fluorescent region with cell-surface CD2, or cortical F-actin at the contact site with the target cell, or having some contact with the glass surface in x-z experiments. The shortest distance from the centroid of the accumulated fluorescence and the cell membrane (identified via differential interference contrast [DIC] microscopy) or the edge of the nucleus (identified via DAPI staining) was also measured using Volocity in some experiments to further define and evaluate polarization. In experiments using siRNA-transfected YTS cells, perforin was visualized using FITC-conjugated mouse anti-human perforin clone 27–35 (BD Biosciences) to identify the effector cell in conjugates formed between YTS and KT86 cells.

**RNA interference and nucleofection.** siRNA containing oligonucleotides with specificity for either CIP4 (Dharmacon siGenome SmartPool) or GAPDH (Dharmacon siGenome Duplex) control were introduced into YTS cells using a nucleofector (program O-017 with reagent R; Amaxa). Titration experiments using CIP4 Western blotting and CIP4 flow cytometry were performed to determine the optimal concentration of CIP4 siRNA, the optimal incubation time after nucleofection. 2.5 μM of siRNA, 2.5–3 × 10⁶ cells per cuvette, and 60 h of incubation after nucleofection were ultimately chosen. Although GAPDH expression could be suppressed using as little as 100 nM siRNA, the use of higher concentrations to control for the CIP4 siRNA did not have any non-specific effect. Constitutively active Cdc42V12 (39), as a GFP–Cdc42V12 fusion construct (provided in expression vector by M. Chou, University of Pennsylvania, Philadelphia, PA) (25), was nucleofected into YTS cells using 2 μg of plasmid, along with either CIP4 or GAPDH siRNA, and incubated for 48 h before use in experiments. 50–60% of cells were GFP positive after incubation. Localization of Cdc42 V12 was determined by microscopy, and increased active Cdc42 was confirmed by an active Cdc42 pull-down assay.

**Statistical analysis.** All graphs denote mean values, except where noted, and error bars represent the SD. Comparisons of groups of data were performed using the two-tailed Student’s t test, and statistical significance is shown in figures (*, P < 0.01, unless otherwise noted).

**Active Cdc42 pull-down assay.** Active Cdc42 was isolated using the Ez-detec Cdc42 activation kit (Pierce Chemical Co.), according to the manufacturer’s protocol. In brief, clarified YTS cell lysates prepared from equal numbers of cells were treated with 0.1 nM GTPγS in the presence of 10 μM EDTA (pH 8) at 30°C for 15 min to activate Cdc42, after which the nucleotide exchange reaction was terminated by adding MgCl₂ and placing samples on ice. Treated lysates were incubated with GST–Pak1–PBD in the presence of SwellGel Immobilized Glutathione (Pierce Chemical Co.) at 4°C for 1 h. After incubation, the mixture was centrifuged at 2,000 g, and the precipitate was washed three times before boiling to isolate bound active Cdc42. Cdc42 was identified in the precipitate by Western blotting using mAb anti-Cdc42.

**Online supplemental material.** Fig. S1 demonstrates the location of the pericentrin–defined MTOC and CIP4 in cytolytic and noncytolytic conjugates in YTS cells. Fig. S2 contains an analysis of colocalization in the 5-μm² region containing the MTOC and an enlargement of detail within that region in cytolytic and noncytolytic conjugates in YTS cells. Fig. S3 depicts perform localization, along with the MTOC, and cytolytic IS in YTS cells overexpressing WT CIP4 or truncated CIP4. Fig. S4 demonstrates CIP4 localization relative to the pericentrin–defined MTOC and IS in eX vivo NK cells. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20061893/DC1.

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**REFERENCES**


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