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To cite this version:

Phosphorylation of Collapsin Response Mediator Protein 2 (CRMP2) on Tyr479 Regulates CXCL12-induced T Lymphocyte Migration

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Running title: CRMP2 transduces chemokine signaling
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In the central nervous system, Collapsin Response Mediator Protein-2 (CRMP2) is a transducer protein that supports the Semaphorin-induced guidance of axons towards their cognate target. However, we previously showed that CRMP2 is also expressed in immune cells and plays a crucial role in T lymphocyte migration. Here we further investigated the molecular mechanisms underlying CRMP2 function in chemokine-directed T-cell motility. Examining Jurkat T-cells treated with the chemokine CXCL12, we found that 1- CXCL12 induces a dynamic re-localization of CRMP2 to uropod, the flexible structure of migrating lymphocyte, and increases its binding to the cytoskeletal protein vimentin; 2- CXCL12 decreases phosphorylation of the GSK-3β targeted residues CRMP2-Thr509/514; 3- tyrosine Y479 is a new phosphorylation CRMP2 residue and a target for the Src-family kinase Yes. Moreover, phospho-Y479 increased under CXCL12 signaling while phospho-Thr509/514 decreased. The functional importance of this tyrosine phosphorylation was demonstrated by Y479-F mutation that strongly reduced CXCL12-mediated T-cell polarization and motility as tested in a transmigration model and on neural tissue. We propose that differential phosphorylation by GSK-3β and Yes modulates the contribution of CRMP2 to cytoskeletal reorganization during chemokine-directed T-cell migration. In addition of providing a novel mechanism for T lymphocyte motility, our findings reveal CRMP2 as a transducer of chemokine signaling.

INTRODUCTION
T-lymphocyte migration is the basis of major immune functions such as responses to infection and inflammation, as well as normal recirculation through the lymphoid organs. Indeed, the role of T-cells depends strongly on their ability to travel between organs via the blood and lymph and to move rapidly within these tissues, by extravasation (1). This latter function is dependent on extracellular signals, among which chemokines play a major role.
Chemokines form a superfamily of small proteins that orchestrate lymphocyte polarization and migration (2). These proteins exert their functions by binding specific seven-transmembrane-domain G-protein coupled receptors on the T-cell surface (3). T-lymphocytes exposed to chemokines, in a soluble or surface-bound gradient, develop a polarized shape, extending at the front, an F-actin-rich lamellipodium, which constitutes the leading edge, and a trailing edge or uropod in which both the microtubule and vimentin networks are retracted during migration. While F-actin has the well-known function of producing the mechanical forces required to generate movement (4), the role of microtubules and vimentin in T-cell migration requires further investigation.
Cytoskeletal remodeling is of key importance in migrating cells (5) and is one of the functions carried out by the chemokine stromal cell-derived factor-1α (SDF-1α), also named CXCL12. In association with its cognate receptor CXCR4, CXCL12 is a potent chemoattractant for mature T-cells and monocytes (6). Following ligand recognition and binding, CXCR4 signaling starts with the activation of G proteins, followed by
various signaling cascade effectors, including MAP kinases, PI3K and PLCγ (7). While this intracellular signaling cascade has not been completely elucidated, the Src family non-receptor tyrosine kinase Lck and the Syk kinase ZAP-70 have emerged as the main candidates for delivering the input signal following CXCR4 activation (8). Thus, tyrosine kinase activity appears as a central step in CXCR4-dependent chemotaxis.

While searching for molecules involved in T-cell motility, we recently identified Collapsin Response Mediator Protein 2 (CRMP2) (9), a protein first described in the context of neuronal growth cone advance (10,11). We demonstrated that CRMP2 regulated both T-cell polarization and spontaneous/chemokine-induced migration of T-lymphocytes. Moreover, CRMP2 was found at the uropod of motile T-cells and have the ability to bind cytoskeletal elements including vimentin. A correlation between CRMP2 expression levels and cell migratory rates towards a chemokine gradient, including CXCL12, was demonstrated by over-expression and knockdown experiments in T-cells (9). In addition, we recently reported that, in mouse model of neuroinflammation, elevated CRMP2 expression in T lymphocytes correlated with their elevated migratory rates and their ability to target the CNS (12). The importance of CXCL12 in the central nervous system (CNS) and its implication in the pathogenesis of CNS disorders, including neuroinflammatory diseases, are well documented (review in (13)). Thus, the aim of the present study was to determine whether and how CRMP2 participates in the transduction pathway induced by CXCL12 on T lymphocytes.

**EXPERIMENTAL PROCEDURES**

**Cells and Antibodies**
The Jurkat T cell line was cultured in RPMI 1640 supplemented with 10% fetal calf serum. Primary T lymphocytes selected from the blood of a healthy donor were cultured for one to two weeks in RPMI complemented with 10% AB human serum and IL2 (20U/mL).

Rabbit polyclonal antibodies recognizing both full-length and cleaved CRMP2 forms have been described previously (14). The peptide sequences used to generate C-ter and pep4 antisera were localized between AA557-572 and AA454-465 in the CRMP2 sequence, respectively. Antibodies were purified by affinity chromatography on the corresponding immobilized peptide. Sheep antisera recognizing CRMP2-pSer522 and CRMP2-pThr509/514 were from KinaseSource Limited (Dundee, UK). Our rabbit polyclonal antibody was raised against a peptide phosphorylated on Y479 (CRMP2-pY479) and purified in a two steps method by affinity chromatography on the corresponding immobilized peptide (step1: elimination of antibody anti un-phosphorylated peptide, step 2: purification of anti phosphorylated peptide). ELISA performed against CRMP2-Y479 and CRMP2-pY479 peptides showed the specificity of our anti-CRMP2-pY479 antibody. In addition, treatment of T-cell lysate with phosphatase CIP significantly reduced the positive signal of anti-Tyr479 but not of anti-pep4 antibody analyzed in WesternBlot (Supplement Figure-1). Rabbit polyclonal antibody anti Yes kinase was from Upstate®. Rabbit polyclonal antibody against the human phospho-Yes1-Y537 was from Abgent® and showed cross-reaction with phospho Src-Y530 a protein with very high homology. Mouse anti Vimentin/LN6 was from Calbiochem®. Anti Erk and phospho-Erk antibodies from Cell Signaling recognized un-phospho- and phospho-p44/p42 MAP Kinase (Erk1 and Erk2). Anti Cdk5, CDK5-pTyr15, Cdk5-pSer159 antibodies were from Santa Cruz Biotechnology. Rabbit anti GSK-3 was from Chemicon International. Mouse anti-pGSK-3 from Upstate Millipore recognized the active forms of GSK-3β (pTyr279) and GSK-3β (pTyr216). Magnetic phospho enrichment beads (TALON® PMAC) were purchased from Clontech. Cell permeable peptide inhibitor of GSK-3β, L803-mts was from Calbiochem®.

**Plasmids and constructs**
The CRMP2-Flag-wt plasmid has been described previously (14). Briefly, full-length CRMP2 was amplified by PCR and inserted directionally into the pCMV2-FLAG vector (Sigma, l’Isle d’Abeau, France). *Mutation:* A two steps PCR procedure was used to generate the CRMP2-Y479F mutant. First, a C-terminal fragment (471-572) containing the Y479-F mutation was generated using a reverse primer introducing an Eco RI site at the 3’ end and a forward primer with the codon: Y 479 (TAC) substituted with F (TTC). Next, this mutated fragment was used as a reverse primer in the second PCR reaction with a wild type forward primer introducing a Hind III site at the 5’ end. The final PCR product was cloned into the Hind III and Eco RI sites of the pCMV2-Flag vector.
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and the DNA sequence of the mutant was verified by sequencing. **Transfection:** Jurkat T cells were transfected with CRMP2-Flag-wt, CRMP2-Flag-Y479F and empty-Flag plasmids using Amaza Nucleofector technology (Köln, Germany), according to the manufacturer’s instructions. T cells were used 18 h after transfection. Transfected cells were visualized by immunostaining with anti-Flag antibody. The percentage of transfection reached 40–50% for most of the Flag constructs.

**Immunocytochemistry**
The CRMP2 forms, Yes kinase and intermediary filament vimentin were detected by indirect immunofluorescence on Jurkat and primary T-cells adhered to collagen I-coated slides (20 μg/ml) and fixed following treatment (acetone – 20°C; 10 min). Cells were incubated with specific antibody (1 h, 37°C) then with Alexa 488- or 546-conjugated anti-mouse or anti-rabbit or anti-sheep IgG antibodies (1h, 37°C) and examined using the Axioplan II fluorescence microscope (Carl Zeiss). Nuclear counter-staining was performed using a fluorescent DNA intercalant, 4', 6'-diamidino-2-phenylindole (DAPI, Boehringer Mannheim).

**Protein interaction assay (pull-down GST-CRMP2)**
One hundred microliters of Jurkat cell lysate, prepared as above, were added either to 80 μl of GST-CRMP2 protein fusion or to 80 μl of GST protein coupled with glutathione-Sepharose 4B (Pharmacia Biotech) (1h at 4°C). GST-CRMP2 and GST beads were washed four times (50 mM Tris (pH 7.4), 1 mM EDTA, 150 mM NaCl, and 0.5% Nonidet P40), and proteins bound to CRMP-2 or to GST beads alone were eluted. GST (in GST beads), Yes and CRMP2 (in GST-CRMP2 beads) and GST (in GST and GST-CRMP2 beads) were revealed by Western blotting.

**Western blotting**
Following CXCL12 treatment, cells were lysed in homogenization buffer (Tris 20mM, EDTA 1mM, EGTA 5mM, sucrose 10%, pH7.4) complemented with phosphatase inhibitors (Na fluoride 5mM, Na pyrophosphate 1mM, β-glycerophosphate 1mM, orthovanadate 1mM) and with protease inhibitor cocktail Complete™ (Roche). Lysates were submitted to ultrasound to dissociate cell aggregates and total proteins measured by Lowry assay (Bio-Rad). Protein samples (10–20 μg) were subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) under reducing conditions and transferred to nitrocellulose membranes (BA85; Schleicher & Schuell Microscience) previously incubated with blocking solution (PBS, 0.1% Tween 20, 5% nonfat dried milk, 1h) and blotted against specific antibody (overnight, 4°C), followed by (1h, room temperature) with rabbit and sheep IgGs antibody coupled with horseradish peroxidase (HRP) and a chemiluminescence (ECL) detection system (Covalab Lyon, France). Densitometric quantification of the immunoblot band was performed using Image Quant (Molecular dynamics) and the data expressed as ratios to the amount detected before any treatment.

**Database and Structure analysis**
The prediction programme PROSITE (http://us.expasy.org/prosite) was used to identify the putative tyrosine kinase site on CRMP2. The structure of CRMP2 was modeled, based on the coordinates available for CRMP2 chain D (15) (protein Data Bank entry 2GSE), using Viewerlite/4.2 (Accelrys).

**Cloning and expression of CRMP2**
The coding sequence for human CRMP-2 (NM_1386) was subcloned into the expression vector pEt21b (Novagen), resulting in a construct with an N-terminal hexahistidine tag. The plasmid was transformed into Escherichia coli BL21(DE3) cells. For expression, cells were grown in 1500 mL Terrific Broth (containing 7% glycerol, 50 μg/mL kanamycin and 100 μL BREOX) in bubble flasks. Cells were grown at +37°C until an optical density of 2.5 at 600 nm was reached. The cultures were cooled to +18°C for 1 h in a water bath. The expression of CRMP-2 was induced by the addition of 0.5 mmol/L IPTG, and expression was allowed to continue overnight at +18°C. Cells were harvested by centrifugation, and the pellets were suspended in lysis buffer (20mM Tris, 500mM NaCl, 1mM DTT, 20% glycerol, 0.1% triton, 10mM imidazole) supplemented with Complete EDTA-free protease inhibitors (Roche, Basel, Switzerland) and 2000 U of benzonase. The solution was sonicated for several cycles on ice. The samples were centrifuged at 14,000 xg for 30 min at 4°C, and the supernatants were incubated with 1.5 ml Ni-NTA resin 50% re-suspended in lysis buffer (QIAGEN) at 4°C for 90 minutes. His-tagged proteins were purified from Ni resin in a wash buffer (20mM Tris, 500mM NaCl, 1mM DTT, 20% glycerol, 0.1% triton, 20mM imidazole) and were eluted with elution buffer (wash buffer + 150 mM imidazole) in 1ml
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fractions. Fractions were evaluated by SDS-PAGE.

Yes in vitro kinase assay
Prior to the assay, His-tagged CRMP2 was dialyzed in buffer (40mM MOPS, 0.5mM EDTA, 5% glycerol) overnight at 4°C using the Float-Allen technology (Interchim), according to manufacturer instructions. For the Yes kinase assay, 0.6 µg of dialyzed His-tagged CRMP2 were incubated with 20 ng of recombinant full-length human Yes (Millipore) diluted beforehand in enzyme dilution buffer (20mM MOPS pH 7, 1mM EDTA, 0.01% Brij, 0.1% β-mercaptoethanol, 5% glycerol). The reaction was allowed in 50 µl of reaction buffer (8mM MOPS, 0.2mM EDTA, 30mM MgCl₂, 2mM EGTA, 10mM β-glycerophosphate, 0.4mM Na₂VO₄, 0.4mM DTT, 200µM ATP) at 30°C for 30 minutes. The reaction was stopped with loading buffer and the mixture was resolved on SDS-PAGE gels.

Transmigration assay
T cell transmigration was performed with Jurkat T cells both in micro-Transwell systems (Costar Transwell Supports -A) and in organotypic cultures of mouse brain (B). A: Transmigration was performed in triplicate in Transwell systems (Boyden chamber, Costar, 5-µm diameter pore size membrane), as described previously (9). Briefly, the T-cell preparation (3 x 10⁶ cells/well) was added in the upper chambers and CXCL12 in the lower compartment (10 ng/mL). Following a 2h incubation at 37°C, cells migrating in the lower chambers were counted under the microscope (at least 30 fields examined). B: T-cell transmigration on neural tissue was assayed on hippocampal cultures prepared as follows. Hippocampi from postnatal (P7) C57BL6 mice were dissected and placed immediately in cold Gey's balanced solution supplemented with glucose (6.5 mg/ml). Four hundred micrometer slices were cut perpendicularly to the septotemporal axis of the hippocampus using a McIlwain tissue chopper. Slices were carefully trimmed for excess tissue, and 6 slices were placed immediately on 30 mm semi-permeable membrane inserts (Millicell-CM, Millipore) in a 6-well plate, each well containing 1 ml of culture medium. The culture medium consisted of 50% Minimum Essential Medium (Gibco), 25% Hank’s balanced salt solution, 25% heat-inactivated horse serum (Gibco), 1% l-glutamine 200 Mm (Gibco) and 6.5 mg/ml D-glucose. Plates were incubated at 37 °C and 5% CO₂. The culture medium was exchanged twice a week. Jurkat T cells (1.10⁶ cells per slice) stained ex vivo using the vital fluorochrome carboxyfluorescein succinimidyl ester/CFSE (1 mM, 5 min, 37 °C) were spotted close to the hippocampus slices (one week culture). Following 18 h contact at 37 °C, slices were extensively washed with D-MEM, fixed with ethanol (10 min, 4 °C) and incubated with DAPI for nuclear counter-staining. The number of infiltrating lymphocytes, counted under fluorescence microscopy, decreased in T-cell populations transfected with CRMP2-Y479-F (graph).

Statistical analysis
Statistical significance in comparing two means was tested with the unpaired Student's t test, p values < 0.05 were considered significant. In the migration test, the number of migratory lymphocytes was counted by light microscopy (15–20 microscope fields per condition - 2 or 3 independent experiments) and data expressed as the mean number of migratory lymphocytes per field.

RESULTS
CXCL12 induces CRMP2 polarization at the T lymphocyte uropod.
To define a link between chemokines and CRMP2, we first examined the localization of CRMP2 in Jurkat T-cells under CXCL12 signaling. We used two different anti-CRMP2 antibodies (anti-C-ter and anti-pep4) that recognize the full-length and cleaved products of CRMP2. An immunofluorescence study of untreated T-cells revealed that CRMP2 was found within the T-cell cytoplasm as punctate dots (Figure 1A, 0mn). Under CXCL12 treatment, CRMP2 moved to the cell trailing edge within 2 minutes and showed quasi-exclusive uropod localization in most polarized cells after 10 minutes treatment (Figure 1A, anti-C-ter antibody). This phenomenon of CRMP2 polarization was still observed after 30 minutes of treatment (not shown). Un-treated Jurkat T-cells showed an asymmetrical CRMP2 distribution in cells, but increases of 1.6 to 2 fold were observed after CXCL12 treatment (Fig 1A, graph). Similar CRMP2 re-localization was observed with anti-pep4 antibody staining (not shown). In addition CRMP2 distribution to the uropod was concomitant with the re-localization of vimentin,
which was quickly redistributed at the trailing edge of polarizing T-cells (Figure 1B, right panel). Interestingly, CRMP2 re-localization was reversed to a great extent (35% decrease) in the presence of AMD3100, an antagonist of the CXCL12 receptor (CXCR4), consequently confirming the specificity of the CXCL12-induced response (Figure 1C). These results supported the idea that chemokines can induce a dynamic re-localization of CRMP2 in T lymphocytes in concert with vimentin, namely in the flexible uropod structure.

**CXCL12 modulates CRMP2 binding to the cytoskeleton.**

It is well known that T-cell uropods are rich in vimentin and microtubules (5), two cytoskeletal elements that have both been described as CRMP2 binding partners (9,16) and actors in T-lymphocyte polarization and migration (17). This led us to hypothesize that CXCL12 could modulate CRMP2 binding to the cytoskeleton to promote T-cell motility. Following CXCL12 treatment (100ng/mL, 10 and 30mn), sub-cellular fractionation was performed on Jurkat T-cell extracts to isolate cytoskeletal elements and associated proteins from the cytosol fraction. Identification of the sub-cellular fractions using antibodies against vimentin, tubulin PARP and Hsp90 indicated that there was no contamination between cytoskeletal and cytosolic fractions (not shown). The cytoskeletal fraction displayed the intermediate filament vimentin but was free from tubulin, probably due to de-polymerization as it is found in the cytosol (not shown). Different fractions were then subjected to Western blotting using anti-CRMP2 antibodies. In whole cell lysates of untreated cells, anti-C-ter antibody revealed CRMP2 bands corresponding to the previously described full-length CRMP2 (62kDa) and bands with higher molecular weight (Figure 2A). Anti-pep4 antibody mainly recognized a 58kDa band, corresponding to the cleaved form of CRMP2, as reported in neural cells (14). After CXCL12 treatment, the efficiency of which was assessed by Erk1/2 phosphorylation (Figure 2A), no difference in CRMP2 expression was detectable in whole cell lysates. However, the distribution of CRMP2 forms differed according to the T-cell compartment examined. Full-length CRMP2 and higher molecular weight bands were found in the cytosolic fractions. These did not show major alterations under CXCL12 treatment. CRMP2 was also found, to a lesser extent, in the cytoskeletal fractions as 62kDa full-length and 58kDa cleaved forms. Interestingly, the expression of both forms was enhanced following CXCL12 treatment. It should be noted that the majority of cleaved CRMP2 was found in the nucleus (not shown), as previously reported in neural cells (14) and was not modified under CXCR4 activation. These results showed that CRMP2 was distributed in the cytoskeletal compartment of T lymphocytes and that CXCL12 had the ability to alter this distribution, enhancing CRMP2 association with cytoskeletal elements.

**CXCL2 increases CRMP2 phosphorylation.**

Functional regulation of CRMP2 in neural cells is mainly dependent on its phosphorylation state, notably via GSK-3β and Cdk5 kinase activity (18). We therefore asked whether, in T lymphocytes, CXCL12 could modify CRMP2 binding to the cytoskeleton through modulation of its phosphorylation. To evaluate CRMP2 phosphorylation, we performed a phosphoprotein enrichment assay (TALON® PMAC, Clonetech) on whole cell extracts of Jurkat T-cells following CXCL12 treatment (100ng/ml) and carried out immunoblotting on the non-phosphorylated (flow through) and phosphorylated (eluate) fractions by Western blotting at 2, 5, 10 and 30 min post-treatment. The full-length CRMP2 forms revealed by the anti-C-ter antibody were present in both the un-phosphorylated and phosphorylated fractions (Figure 2B). In contrast, the cleaved form of CRMP2 was only found in the phosphorylated protein pool, indicating that this form is mostly phosphorylated. CXCL12 treatment rapidly increased the level of CRMP2 phosphorylated forms, peaking at 2 min post treatment and still high at 30 min. The efficiency of the phosphoprotein enrichment procedure was ascertained by phospho-Erk1/2 immunoblotting, which confirmed the specific presence of phosphorylated proteins in the eluate and at the same time, the increase following CXCL12 treatment. Similar experiments performed on primary T-lymphocytes isolated from healthy donors showed similar observations (not shown). A more precise evaluation of CRMP2 phosphorylation in response to CXCL12 was carried out using anti-CRMP2-pSer522 and anti-CRMP2-pThr509/514 antibodies recognizing two sites targeted by Cdk5 and GSK-3 kinases, respectively (Figure 3A). Immunoblotting of Jurkat cell lysates showed that CRMP2-pSer522 and CRMP2-pThr509/514 were present as full length 62kDa CRMP2 in T-cells and were variably expressed during chemokine treatment.
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the efficiency of which was ascertained by phosho-Erk1/2 detection. While Ser522 phosphorylation was found at relatively low levels, Thr509/514 phosphorylation decreased quickly by 4 min and was undetectable thereafter. This was consistent with the activity of Cdk5 and GSK-3 kinases evaluated by the detection of Cdk5-pTyr15, Cdk5-pSer159, GSK-3α-pTyr279 and GSK-3β-pTyr216, the active forms of these kinases (Figure 3B). Cdk5 displayed a stable level of phosphorylation on Tyr15 and Ser159, reflecting a conserved level of Cdk5 activation. In contrast, GSK-3 exhibited de-phosphorylation mainly detected on the GSK-3β isoform, revealing a decreased activity starting at 4 min post-treatment. Taken together, these results first revealed, as previously described in neural cells, that the CRMP2 residues Ser522 and Thr509/514 could be phosphorylated in T lymphocytes. More importantly, they demonstrate that CXCL12 triggers a signaling cascade leading to differential modulation of CRMP2 phosphorylation of these residues, namely with a net decreased phosphorylation on Thr509/514. Intriguingly, these modulations were mainly detected on the full-length CRMP2 forms, while phosphoprotein enrichment assays (Figure 2B) showed a strong phosphorylation of the cleaved CRMP2 form following chemokine treatment. This led us to suspect the participation of an additional phosphorylated target in the response of CRMP2 to CXCL12.

Tyrosine 479 is a new phosphorylation residue in CRMP2 sequence

It is known that CXCL12 triggers a tyrosine phosphorylation cascade in T-lymphocytes, which involves the serial recruitment and activation of tyrosine kinases including Lck, ZAP-70 and Itk (8). We therefore searched for tyrosine target residues potentially modulated under chemokine treatment by analyzing CRMP2 protein sequences. A database study of the 572 amino acids identified tyrosine 479 (Y479) as a potential new phosphorylation residue, located in the phosphorytosine consensus motif KxxxDxxY within residues 472-479 (Figure 4A). In addition, inspection of this region also showed the presence, close to Y479, of a putative SH3-binding motif of the form RxxPxP within residues 467-473. In order to assess the accessibility of these sequences to binding protein partners, we evaluated the position of both Y479 and the SH3-binding motif within the known structure of CRMP2 (Figure 4A) based on the coordinates available for fragment 15-489 (15). Surface exposure representation of this CRMP2 form revealed that, in contrast to Y479, the putative SH3-binding motif was exposed, suggesting a possible binding with SH3-domain bearing proteins (Figure 4A insert). It has been shown that interaction with the SH3 domain-binding motif induces protein conformational changes (19), so the latter could be the basis of subsequent Y479 exposure. These observations suggested Y479 as a major putative phosphorylation tyrosine within the CRMP2 sequence.

CRMP2 tyrosine-phosphorylation is carried out by the Src-family kinase Yes and increases under chemokine treatment.

In view of the presence of the putative SH3-binding motif close to the potential phosphorylatable site Y479, a possible interaction between CRMP2 and tyrosine kinases through its SH3 domain was studied. This was done using a membrane array bearing several protein SH3 domains that remain folded in active conformations. Ten different lymphocyte tyrosine kinases, including Abelson kinase (Abi), Src family kinases (Lck, Yes, c-Src, Fyn, Hck, Blk) and Tec family kinases (Itk, Txk, Btk) were present in this array. Following His-tagged recombinant CRMP2 hybridization, protein-protein interactions were visualized using anti-His antibody and spot intensities revealed the interaction strength (Table I). Yes kinase displayed a strong binding to CRMP2, while Blk and Abi showed weak signals. In addition, four non-kinase protein SH3-domains belonging to Vav1, PLCγ, ITSN and PI3β also displayed strong binding to CRMP2. Interestingly, PLCγ and ITSN have previously been observed as binding partners for CRMP2 (20,21). We then focused on Yes, the more potent tyrosine kinase candidate for CRMP2 phosphorylation.

The Yes /CRMP2 interaction was evaluated by several approaches. First, localization of these proteins was assessed on primary T lymphocytes and Jurkat T-cells that had been allowed to adhere onto collagen-I coated coverslips and then treated with CXCL12 (100ng/ml, 5 min) (Figure 4B). Immunofluorescence, performed with anti-Yes and anti-ep4 antibodies, showed the co-distribution of CRMP2 and Yes, especially at the uropod of polarized T-cells. Yes /CRMP2 interactions were next examined by a GST-pulldown assay using cell lysates from primary T-lymphocytes and from neural cells.
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(Dev cell line) (Figure 4C), as CRMP2 is also involved in motility in the central nervous system (CNS). CRMP2 immobilized on glutathione-Sepharose beads was incubated with cell lysates. Western blots, performed on eluates from both cell types, showed the presence of Yes protein in association with CRMP2-GST, but not with GST alone. Taken together, these results defined the Yes kinase as a potent binding partner for CRMP2. In order to evaluate the functional significance of this interaction, an in vitro kinase assay was performed using active recombinant human Yes kinase and His-tagged CRMP2 as a substrate (Figure 4D). Phosphorylation was detected using an anti-phospho-Tyrosine antibody by immunoblotting. A control was carried out in the absence of CRMP2, which showed Yes self-phosphorylation. A band corresponding to CRMP2 phosphorylation was detected only in the presence of ATP. As a consequence of protein phosphorylation, this band displayed a slight increase in molecular weight.

To confirm the presence of tyrosine-phosphorylated forms of CRMP2 in T-cells, a polyclonal antibody was raised against a fragment of the CRMP2 sequence containing the phosphorylated residue Tyr479. Immunoblotting with this antibody revealed the presence of CRMP2-pTyr479 in T-lymphocytes, detected as both full-length and 58-kDa cleaved proteins. Examination of CRMP2-pTyr479 in Jurkat T-cells treated with CXCL12 (100ng/ml; 0, 2, 5, 8, 14, 20, 30 min) showed an increase in Tyr479 phosphorylation (Figure 5A), mainly observed at 8-14 min post-treatment. Similar data were observed in 5 independent experiments. Increased expression of CRMP2-pTyr479 was concomitant with the decrease of CRMP2-Thr509/514 as observed in Figure 3A. This suggested a relationship between the phosphorylated Thr509/514 and Tyr479 sites. This hypothesis was evaluated by blocking GSK-3β activity using a cell permeable peptide inhibitor (L803-mts, Calbiochem, 20, 40 mM, 4h). For this purpose we used a virus-infected T-cell line (C91PL) that constitutively expressed CRMP2-pTyr479, possibly due to chronic activation. As expected, GSK-3β blockade resulted in the decrease in CRMP2-pThr509/514. In parallel, CRMP2-pTyr479 increased, validating our hypothesis (Supplemental Figure 2). Immunoblotting using antibody directed against a phospho-Yes1-Y537 peptide revealed the constitutive presence of p-Yes in the Jurkat T-cells, that was not significantly modified by CXCL12 treatment. This indicated that CRMP2-Y479 phosphorylation by Yes under CXCL12 signaling might occur in Jurkat T-cells but mainly depended on site accessibility to this kinase.

Immunofluorescence was then performed on Jurkat cells treated with CXCL12 (15 min) using antibodies against the phosphorylated and non-phosphorylated CRMP2 forms. Staining for CRMP2-pTyr479 was mainly with a polarized distribution in T lymphocytes (Figure 5B). Co-localization with vimentin showed that, compared to the CRMP2 forms recognized by the anti-pep4 and anti-C-ter antibodies, which were either associated (orange as merged staining) or not associated (green staining) with vimentin, respectively, the phosphorylated CRMP2-Tyr479 was mainly colocalized with vimentin at the trailing pole (Figure 5 B arrows). Taken together, these results identified a new form of phosphorylated CRMP2 that was modulated by CXCL12 signaling, colocalized with cytoskeletal elements and could be targeted by the Src-family kinase Yes.

CRMP2-Tyr479 phosphorylation is involved in chemokine-induced polarization and migration of T-cells.

To assess the functional significance of CRMP2 phosphorylation on Tyr479, we engineered the mutation Y479-F on the full-length CRMP2 sequence. The effect of Tyr479 phosphorylation impairment on T-cell polarization was then analyzed in Jurkat T-cells transiently transfected with Flag-tagged CRMP2-wt and CRMP2-Y479-F mutant. Twenty-four hours after transfection, T-cells were allowed to adhere onto collagen-I-coated slides, then treated with CXCL12 and examined by fluorescence microscopy. Polarization of CRMP2 in transfected T-cells, as visualized by Flag-positive immunostaining (Figure 6- A, D, G, J), was examined based on vimentin network co-localization (Figure 6- C, F, I, L, orange as merge staining). This allowed us to evaluate the polarization of Flag positive T-cells transfected either with CRMP2-wt or CRMP2-Y479-F (expressed as a percentage of all transfected cells). The un-treated Jurkat T-cell population transfected with CRMP2-wt displayed ~28% spontaneously polarized cells, but this clearly decreased in T-cells transfected with the CRMP2-Y479-F mutant (Figure 6- graph). Following CXCL12 treatment, vimentin was quickly redistributed to the uropod in CRMP2-wt transfected T-cells (Figure 6- H, I). In contrast,
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CRMP2-Y479-F transfected cells were clearly less polarized (Figure 6- K, L and graph). In addition, the increase in the number of polarized cells following CXCL12 treatment was lower in Jurkat T-cells transfected with CRMP2-Y479-F than with CRMP2-wt (31% versus 42%, respectively) (Figure 6, graph), thus confirming the impact of Tyr479 phosphorylation on T-cell polarization. These results clearly showed the role of CRMP2-Y479 phosphorylation in T lymphocyte polarization.

As T-cell polarization is a prerequisite for migration, we further evaluated the influence of Tyr479 phosphorylation on T-cell migration. Thus, we first assessed the ability of transfected Jurkat T-cells to migrate towards CXCL12, by performing a transmigration assay in Transwell chambers. As shown in Figure 7A, the rate of migration of CRMP2-Y479-F transfected cells was drastically reduced compared to those with CRMP2-wt and control cells (empty vector). Beyond T-cell transmigration that is necessary to traverse blood vessels, migration within invaded tissue is also a key point, especially within the CNS where CXCL12 and its cognate receptor are constitutively expressed. We therefore examined whether Tyr479 phosphorylation had an influence on T-cell migration within neural tissue, using mouse hippocampal organotypic culture. Transfected Jurkat T-cells (40-50% transfection efficiency) were stained with the vital dye CFSE in order to easily visualize them both on and in neural tissue (Figure 7B). Cells were then spotted close to brain slices and were counted after 18 hours incubation. CRMP2-Y479-F transfected cells displayed a reduced ability to travel on neural tissue compared to wild type transfected cells. These results demonstrated the role of CRMP2-Y479 phosphorylation in the process of T-cell migration within neural tissue.

DISCUSSION

To date, CRMP2 has essentially been known to specify neuronal polarity and promote axon elongation and branching in response to semaphorin by regulating microtubule assembly, reorganizing actin filaments and protein trafficking (22,23). We recently showed that CRMP2 is expressed efficiently in T lymphocytes and demonstrated that these cells need CRMP2 to establish a polarized morphology and to engage in subsequent chemokine-directed migration (9), defining CRMP2 as a major player in cell polarity and motility within a larger field than just the nervous system. The present study undoubtedly establishes a link between CXCL12/CXCR4 signaling and CRMP2 in T lymphocytes.

CRMP2 functions in chemokine signaling.

Similar to other chemokines, CXCL12, when binding to its receptor CXCR4, activates a series of down-stream signaling pathways including heterotrimeric G-proteins, the Src family tyrosine kinases and phosphoinositide-3-kinase (PI3K), leading to cell polarization (24). The present observations prompted us to propose CRMP2 as part of this uropod signaling pathway. First, we showed that T lymphocytes treated with CXCL12 induced a re-localization of CRMP2 to the uropod of polarizing T-cells, a feature impaired by treatment with the CXCR4-antagonist AMD3100. This chemokine-induced change in intracellular distribution paralleled a modification of CRMP2 phosphorylation, together with an increased ability to bind cytoskeletal components. In addition, several SH3-binding proteins we identified as potential molecular partners of CRMP2 belong to the signaling cascade downstream of CXCR4. Notably, the association of CRMP2 with Vav-1, a GEF with a pivotal role in migratory signal transmission through CXCR4 (25), suggests that CRMP2 is involved in recruitment at the uropod of signaling intermediates that transmit the chemotactic signal for cytoskeletal re-organization.

CRMP2 is involved in cytoskeletal re-organization under chemokine signaling.

Similar to its crucial role as a regulator of cytoskeletal assembly in neural cells in response to semaphorin (16,26,27), CRMP2 is probably involved in the regulation of lymphocyte polarity in response to chemokine. We showed that CXCL12 enhanced CRMP2 recruitment on cytoskeletal elements, notably on vimentin a molecule known to rapidly collapse into an aggregate within the uropod under chemokine signaling (28). This supports the idea of CRMP2 involvement in the maintenance and modulation of chemokine-induced T lymphocyte polarity and migration. As a potent molecular partner of actin, tubulin, kinesin-1 and of the endo-exocytic proteins numb and intersectin (29,30), CRMP2 could contribute to the dynamic reorientation of critical surface receptors/signaling proteins, selective vesicular traffic and contraction/ retraction from the front to the back required for cell movement, all orchestrated by the polarized.
acto-myosin and microtubule cytoskeletons [review in (17)].

**CRMP2 is differentially phosphorylated under CXCL12 signaling.** The exact implication of CRMP2 in chemokine-directed T cell polarization and motility remains to be determined. However this certainly involves subtle changes in CRMP2 phosphorylation state as sequential phosphorylation near the C-terminus by several serine/threonine kinases has been shown to be crucial for CRMP2 function in neural cells (18,31). Briefly, Rho-kinase phosphorylates CRMP2 at Thr555 under ephrin-A5 and myelin-associated glycoprotein signaling (32,33), while Cdk5 phosphorylates CRMP2 at Ser522 in response to Sema3A, priming the protein for subsequent phosphorylation by GSK-3β Thr509 and Thr514 (34). All these post-translational modifications result in a reduced affinity of CRMP2 for tubulin, leading to growth cone collapse and the inhibition of neurite outgrowth. Similar to what has been observed in neural cells, we found that CRMP2 was phosphorylated at Ser522 and Thr509/514 residues in resting T lymphocytes. CXCL12 differentially modulated CRMP2 phosphorylation at those sites, inducing a rapid de-phosphorylation at the GSK-3 target residue Thr509/514, while phosphorylation of the Cdk5 site, Ser522, remained relatively stable. Interestingly, Cdk5 and GSK-3β kinase activity paralleled these phosphorylation states. The major difference in the stability of phosphorylated Ser522 and Thr509/514 sites, recently reported in neural cells (35), could further explain the differential modulation of CRMP2 phosphorylation observed in T lymphocytes in response to chemokine. Indeed, these authors showed that protein phosphatase-1 rapidly de-phosphorylated CRMP2 at Thr509/514 upon GSK-3 inhibition, while the Cdk5 site Ser522 remained resistant to phosphatase treatment. In neural cells stimulated by IGF-1, BDNF and NT-3, GSK-3 inhibition and phosphatase-1 activity on CRMP2 promote CRMP2 de-phosphorylation, supporting microtubule assembly and axon outgrowth (31,34). Accordingly, our present data suggest CXCL12 as a further regulator of CRMP2 phosphorylation at the GSK-3 target site.

Another novelty of our work is the identification of a new phosphorylation site, Tyr479, located at the C-terminus near a putative SH3-binding motif. We showed that Yes, a non-receptor tyrosine kinase of the Src family, has the ability to phosphorylate CRMP2. This feature was compatible with the demonstration of Yes binding to CRMP2 through its SH3-domain. It is worth noting that CRMP2 showed the ability to bind Yes but not Fyn, Hck, Lck , Src and Blk, which belong to the same kinase family. In fact, although Src family members share significant sequence and structural homology, these have been shown to serve not only redundant but also distinct functions (36). Phosphorylation of CRMP2-Tyr479 by Yes kinase under chemokine signaling is conceivable, at least in Jurkat cells as they constitutively expressed the active Yes phosphorylated form and CRMP2-pTyr479 level expression was noticeably enhanced following CXCL12 treatment.

The role of phosphorylation at Tyr479 in functionally controlling CRMP2 under CXCL12 signaling was clear as mutation of Tyr479 profoundly altered CXCL12-induced polarization and greatly reduced the transmigration rate of T-cells and their ability to navigate ex vivo on neural tissue. It is therefore likely that phosphorylated CRMP2 forms recruited at the uropod, notably CRMP2-pTyr479, would be included in the signalosome formed under CXCR4 activation. Intriguingly, CRMP2-pTyr479 later appeared in chemokine-treated lymphocytes. One can propose several explanations for this. Accessibility to the tyrosine kinase site could be a limiting factor. We observed that Tyr479 was not exposed on the surface of the CRMP2 molecule and probably required structural changes to become accessible to SH3-bearing molecular partners such as Yes. In fact, phosphorylation at this tyrosine site appeared when phosphorylation of the GSK-3 target (Thr509/514) was no longer detectable. Furthermore, blockade of GSK-3β activity significantly reduced the pThr509/614 form but increased pTyr479. Thus, de-phosphorylation at Thr509/514 could alter the overall charge of the protein sequence in the vicinity of Tyr479, thus enhance the stability of phosphorylation at that site, in a manner similar to that for the Ser522 site (35).

**The association of CRMP2 with CXCL12 signaling has functional significance.** Given the ubiquitous pairing of CRMP2 and CXCR4, the CXCR4/Src/GSK-3/CRMP2 pathway described here might have consequences in several physiological and pathological situations. In the CNS, CXCL12 functions in localizing immune cells to the perivascular space, thereby limiting the parenchymal infiltration of autoreactive T-cells. However in patients with
CRMP2 transduces chemokine signaling

neuroinflammatory diseases, including multiple sclerosis (MS), CXCL12 localization switches from a basolateral distribution towards the luminal and parenchymal sides of endothelial cells (37). Thus, CXCL12 accessibility at the blood-brain-barrier and the elevated expression found in reactive astrocytes within active and chronic lesions in MS, probably promote T-cell infiltration in brain parenchyma (38). It is therefore tempting to speculate that CRMP2 is involved in the interplay between T-lymphocytes, brain microvasculature and neural cells. Indeed, we have detected elevated CRMP2 expression in activated T-lymphocytes of patients suffering from neuroinflammatory diseases (9) and in the T-cells of mice developing either virus-induced neuroinflammation (12) or experimental autoimmune encephalitis (EAE) [unpublished data]. CRMP2 over-expression in peripheral T-cells was associated with high motility of selected human and mice T lymphocytes and with brain T-cell infiltration and high clinical scores in mouse models. Consequently, elevated chemokine signaling at the BBB and CRMP2 activity in activated T-cell effectors might be required to orchestrate the infiltration of auto-reactive T-cells in brain and thus disease progression in neuroinflammatory diseases.

It also appears that chemokines and their receptors have numerous roles to play in the developing and adult nervous systems (13). CRMP2 could participate in axon guidance and path finding regulated by the CXCL12/CXCR4 axis, as this has been recognized as decisive in regulating the migration of adult neural progenitors and axon projection of spinal motor neurons during development (39). In addition, a promising field of research has opened up in neurooncology as both CXCL12/CXCR4 and CRMP2 are upregulated and hyperphosphorylated in brain tumors (40) (41).

In conclusion, we propose a novel regulation of CRMP2 activity that may be important for both the immune and nervous systems. Further studies should clarify the importance of the CXCR4/CRMP2 pathway either as a potential peripheral biomarker of neuroinflammation and tumor progression or as a target for further development of therapeutic strategies.
REFERENCES

FOOTNOTES
Abbreviations used:  Akt: v-akt murine thymoma viral oncogene homolog; CRMP2: collapsin response mediator protein 2; Cdk5: cyclin-dependent kinase 5; GSK-3: glycogen synthase kinase 3; GPCR: G protein-coupled receptor; MS: multiple sclerosis; PI3K: phosphoinositide-3 kinase; BDNF: brain-derived neurotrophic factor.

ACKNOWLEDGMENTS
This work was supported by grants and fellowships from the French Agency of Research on Multiple Sclerosis (ARSEP), INSERM and Agence Nationale de la Recherche (ANR). The authors thank Stéphan Menigoz for the illustration of the CRMP2 structure and Serge Nataf for critical reading of this manuscript.

FIGURE LEGENDS & TABLE
Table I - Identification of SH3 protein-CRMP2 interaction
CRMP2-His recombinant protein was incubated with a membrane spotted in duplicate with SH3 domains of 38 proteins (TranSignal TM SH3 Domain array I – Panomics) according to manufacturer’s instructions. Anti-His antibody revealed the association of CRMP2 with multiple SH3 domains, including those of some tyrosine kinase proteins. Spot intensities (- to ++++) indicated the binding affinity of SH3 domains to the ligand CRMP2 and revealed Yes as a potent tyrosine kinase candidate for CRMP2

Figure 1: CXCL12 induced a polar distribution of CRMP2 at the T-cell uropod
Jurkat T-cells were allowed to adhere to collagen1-coated slides, treated with CXCL12 (100ng/mL) and examined for CRMP2 distribution at 0, 2, 5 and 10 min post-treatment following fixation. (A) Immunostaining with anti-CRMP2-C-ter antibody (left panel) showed a markedly elevated number of cells with polarized CRMP2 (graphical representation) under CXCL12 treatment. (B) Co-distribution of immunodetected CRMP2 and vimentin at the uropod in CXCL12 treated T-cells. Nuclei stained blue by DAPI (CRMP2 staining using two different specific antibody). (C) Treatment of Jurkat T-cells with the CXCR4 antagonist AMD3100 specifically reduced the number of cells exhibiting polarized CRMP2 under CXCL12 treatment, confirming the association between CXCL12 signaling and CRMP2.

Figure 2: CXCL12 promoted CRMP2 association with the cytoskeleton and enhanced CRMP2 phosphorylation
A- Jurkat cells were treated with CXCL12 (100 ng/mL, 10, 30 min) and cell lysates subjected to subcellular fractionation, then Western blotting using anti-CRMP2-pep4 and -C-ter antibodies. Full-length (open arrow) and cleaved (black arrow) CRMP2 forms were detected in the cytoskeletal fraction (identified by vimentin detection), and increased under CXCL12 treatment (see relative values). The full-length forms distributed in the cytosol, did not show major modification. CXCR4 activation was assessed by increases in phospho-Erk1/2 products (representative experiment from three different experiments). B- Jurkat T-cells were treated with CXCL12 (100ng/mL) and cell lysates subjected to phosphorylated form enrichment at 0, 2, 5, 10 and 30 min post-treatment. Whole cell extract, un-phosphorylated (flow through) and phosphorylated proteins (eluate, authenticated by phospho-Erk1/2 detection) were Western blotted using CRMP2 antibodies. Full-length forms (open arrow) distributed in both phospho and un-phosphorylated fractions, while the cleaved form of CRMP2 (black arrow) was found mainly in the phosphorylated pool. Phosphorylated CRMP2 increased under CXCL12 treatment. CXCR4 activation was assessed by the induction of phospho-Erk1/2 products.

Figure 3: CXCL12 differentially modulated CRMP2 phosphorylation on Cdk5 and GSK-3 target residues.
A- Jurkat T-cells were treated with CXCL12 (100 ng/mL, 0, 1, 2, 4, 6, 10, 15, 30 min). Cell lysate blotting against anti-C-ter CRMP2 and the forms phosphorylated on Ser522 and Thr509/514, targets of Cdk5 and GSK-3β kinases, respectively, showed a decrease in CRMP2- pThr509/514 under
CXCL12 treatment, which was assessed by the induction of phospho-Erk1/2 products. 

**B-** Blotting against the kinases Cdk5 and GSK-3 and against their active forms Cdk5-pTyr15, Cdk5-pSer159, GSK-3α-pTyr279 and GSK-3β-pTyr276 showed the presence and activity of GSK-3 and Cdk5. CXCL12 induced a progressive decrease in GSK-3β-pTyr276 (see relative values).

**Figure 4:** Yes of the Src family is a tyrosine kinase partner for CRMP2

A- Identification of a potential tyrosine phosphorylation residue, Y479, in the phosphotyrosine consensus motif KxxxDxxY (AA 472-479) on CRMP2, close to a putative SH3-binding motif (RxxPxxP). The surface exposure of the SH3 binding motif in the CRMP2 structure is shown in a ribbon diagram revealing the structure of short CRMP2 from residues 15 to 489. The helices are distinguished by their red color, the strands are in green, and the turns are in white. The putative SH3 binding domain involves residues R467, P470, and P473, shown as ball-and-stick representations. These residues are located in a flexible part N-terminal to the last α helix. The inset shows the surface representation of short CRMP2, indicating the surface exposure of residues R467, P470 and P473 (magenta).

B- Co-detection of Yes and CRMP2 in primary T cells and the Jurkat T cell line allowed to adhere to collagen I-coated slides, then treated with CXCL12 (100ng/mL, 15min, orange as merge color). C- Protein interaction assay (GST pull down): sepharose-4B beads coupled to the fusion protein CRMP2-GST and GST were added to the cell lysates of primary T lymphocytes and Dev neural cells. Blotting eluates against Yes, CRMP2 and GST validated the binding of Yes with CRMP2-GST beads and not with GST alone.

D- *In vitro* kinase assay was performed with active recombinant Yes (Millipore) and recombinant CRMP2 protein Western blotting against phospho-Tyr residues detected CRMP2 phosphorylation on tyrosine in the presence, but not in the absence, of ATP. Blotting to CRMP2 detected a slight increase in molecular weight following CRMP2 phosphorylation.

**Figure 5:** CRMP2-pTyr479 is enhanced and polarized in T cells under CXCL12 treatment

A- Detection of CRMP2 phosphorylated on Tyr479 in Jurkat cells with our polyclonal antibody raised against a peptide comprising the phosphorylated Tyr479 residue. Western blotting revealed CRMP2-pTyr479 in full-length and cleaved forms, which increased at 8-14 min following CXCL12 treatment (see relative values). Blotting against phospho-Yes1-Y537 revealed the constitutively active forms of Yes. B- Co-immunodetection of vimentin (red) and CRMP2 (green) in Jurkat cells treated with CXCL12 (100ng/mL, 15min) using anti-CRMP2-pep4 and -C-ter antibodies and against CRMP2-pTyr479. Orange spots (merged color) represent co-localization of CRMP2 forms and intermediate filament vimentin at uropod.

**Figure 6:** CRMP2-Tyr479 is involved in the chemokine-induced polarization of T-cells

Jurkat cells were transfected with either CRMP2flag-wt (A-C; G-L) or CRMP2flag-Y479-F mutants (D-F; J-L), allowed to adhere to collagen I-coated slides then treated with CXCL12 (G-L, 100ng/mL). Polarization of transfected T-cells was visualized by co-detection of flag and vimentin. The quantification of polarized cells, expressed as percentages of all transfected cells, is shown graphically. Mutation of the tyrosine residue (CRMP2-Y479-F mutant) specifically reduced spontaneous and chemokine-induced polarization (graphical representation).

**Figure 7:** Mutation of CRMP2-Tyr479 resulted in reduced T lymphocyte migratory rates

Jurkat cells transfected with either flag vector, CRMP2flag-wt or CRMP2flag-Y479-F plasmids were examined for transmigration towards CXCL12 in Transwell chambers. Mutation of the tyrosine residue specifically reduced the migratory rate (graphical representation). B- For migration analysis on neural tissue, transfected Jurkat cells were stained with the vital dye CFSE then spotted close to hippocampal organotypic slices and examined for migration using fluorescence microscope (green color). Counter-staining with DAPI visualized neural cell nuclei (blue color). Mutation of the tyrosine residue (CRMP2-Y479-F) specifically reduced the ability of T-cells to navigate on neural tissue, as shown by quantification of migrating cells (graphical representation).
**Table I**

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<tr>
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<tr>
<td>Abl Abelson tyrosine kinase</td>
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</tr>
<tr>
<td>BLK B-lymphocyte specific protein tyrosine kinase</td>
<td>+</td>
</tr>
<tr>
<td>LCK Human T-lymphocyte specific protein tyrosine</td>
<td>-</td>
</tr>
<tr>
<td>FYN Proto-oncogen tyrosine protein kinase</td>
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</tr>
<tr>
<td>BTK Bruton Tyrosine kinase</td>
<td>-</td>
</tr>
<tr>
<td>c-Src Cellular rous sarcoma virus oncogen homolog 1</td>
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<tr>
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<td>TXK Tyrosine-protein kinase TXK</td>
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<td>Itk Interleukin-2-inducible T-cell kinase</td>
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CRMP2 transduces chemokine signaling

Figure 1
Figure 2

A

CXCL12

whole cell

10 30 min

- + - +

CRMP2-C-ter

60 60

- + - +

CRMP2-pep4

60 60

- + - +

p Erk1/2 Erk1/2

Cytoskeleton

10 30 min

- + - +

Relative values

1 1.7 1 9.4

B

whole cell

CXCL12

0 2 5 10 30 min

CRMP2-C-ter

80 cr

- + - +

CRMP2-pep4

80 cr

- + - +

GAPDH

86

Flow-through

0 2 5 10 30 min

Eluate

0 2 5 10 30 min

p Erk 1/2 Erk 1/2

vimentin
CRMP2 transduces chemokine signaling

Figure 3

A

CXCL12

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* relative values

B

CXCL12

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* relative values
CRMP2 transduces chemokine signaling

**Figure 4**

**A**

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KVLGEDGLHYEGSCRPRKPPFDEVY 472
KRIKARSRLAEINS 484
KVLGEDGLHYEGSCRPRKPPFDEVY 472
KRIKARSRLAEINS 484
VRNLHQSFLSLGAQIDNIPRT 573
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```
phospho Tyr 80
CRMP2 60
```

19
Figure 5

A

CXCL12

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B

Vimentin

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CRMP2 transduces chemokine signaling
Figure 6

CRMP2 transduces chemokine signaling

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CXCL12

CRMP2 WT

CRMP2-Y479-F

% Polarization / transfected T-cells

CRMP2-wt
CRMP2-Y479-F

+ CXCL12

***

**

+ 31%

+ 42%

+ 42%
CRMP2 transduces chemokine signaling

Figure 7

A

A bar graph showing the number of migratory cells with error bars.

- **CRMP2-wt**
- CRMP2-Y479-F
- empty vector

B

A histogram and images showing the number of migratory cells.

- **CRMP2 wt**
- CRMP2-Y479-F

Images of hippocampus slices with labeled cells.