

Re-visiting the *trans* insertion model for complexin clamping

Shyam S Krishnakumar¹*[†], Feng Li^{1†}, Jeff Coleman¹, Curtis M Schauder¹, Daniel Kümmel^{1,2}, Frederic Pincet^{1,3}, James E Rothman¹, Karin M Reinisch¹*

¹Department of Cell Biology, Yale University School of Medicine, New Haven, United States; ²School of Biology/Chemistry, Universität Osnabrück, Osnabrück, Germany; ³Laboratoire de Physique Statistique, UMR CNRS 8550 Associée aux Unive, Ecole Normale Supérieure, Paris, France

Abstract We have previously proposed that complexin cross-links multiple pre-fusion SNARE complexes via a *trans* interaction to function as a clamp on SNARE-mediated neurotransmitter release. A recent NMR study was unable to detect the *trans* clamping interaction of complexin and therefore questioned the previous interpretation of the fluorescence resonance energy transfer and isothermal titration calorimetry data on which the *trans* clamping model was originally based. Here we present new biochemical data that underscore the validity of our previous interpretation and the continued relevancy of the *trans* insertion model for complexin clamping. DOI: 10.7554/eLife.04463.001

Introduction

The tightly regulated release of neurotransmitters is key to all information processing in the neural circuitry. The fusion of a synaptic vesicle to release the neurotransmitters is mediated by the SNARE (Soluble N-ethylmaleimide-sensitive factor Attachment protein REceptor) complex, which forms between vesicle and target membranes as v-SNAREs emanating from transport vesicles assemble with t-SNAREs emanating from target membranes (*Sollner et al., 1993; Weber et al., 1998; Jahn and Scheller, 2006*). Key proteins regulating SNARE-mediated fusion at the synapse are the calcium sensor synaptotagmin and complexin (CPX) (*Brose et al., 1992; McMahon et al., 1995; Fernandez-Chacon et al., 2001; Giraudo et al., 2006; Sudhof and Rothman, 2009*). Genetic and physiological studies in a number of model systems show that CPX inhibits the spontaneous release of neurotransmitters and is also essential for synchronous exocytosis (*Huntwork and Littleton, 2007; Maximov et al., 2009; Yang et al., 2010; Martin et al., 2011; Cho et al., 2014*). CPX 'clamps' the SNARE assembly process to prevent the continuous release of neurotransmitters (*Giraudo et al., 2010; Martin et al., 2011; Cho et al., 2014*). CPX 'clamps' the snake assembly process to prevent the continuous release of neurotransmitters (*Giraudo et al., 2010; Martin et al., 2011; Cho et al., 2014*). CPX 'clamps' the snake assembly process to prevent the continuous release of neurotransmitters (*Giraudo et al., 2006*). It does so by stabilizing the SNAREs in an otherwise unavailable 'intermediate' energetic state in which the four helix bundle is about 50% zippered (*Li et al., 2011*).

Based on the X-ray crystal structure of CPX bound to a mimetic of this half-zippered intermediate in which only the N-terminal portion (residues 26–60) of v-SNARE, VAMP2, is present (SNARE Δ 60), we proposed a molecular model for the clamping of the SNARE assembly by CPX (*Kümmel et al., 2011*). We found that the CPX central helix (CPX_{cen}, the SNARE-binding domain) binds one SNAREpin while the accessory helix (CPX_{acc}, the clamping domain) extends away and bridges to a second SNAREpin. The CPX_{acc} interacts with the t-SNARE in the second SNAREpin, occupying the v-SNARE binding site, thus inhibiting the full assembly of the SNARE complex. Further, the intermolecular *trans* clamping interaction of CPX organizes the SNAREpins into a 'zig-zag' topology that is incompatible with opening a fusion pore (*Krishnakumar et al., 2011*; *Kümmel et al., 2011*).

We used isothermal titration calorimetry (ITC) to characterize the interaction of the CPX_{acc} with the t-SNARE, fluorescence resonance energy transfer (FRET) analysis to establish the angled conformation

*For correspondence: shyam. krishnakumar@yale.edu (SSK); karin.reinisch@yale.edu (KMR)

[†]These authors contributed equally to this work

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eLife digest Molecules called neurotransmitters are used to carry signals between neurons. The neurotransmitters in the first neuron are stored in small bubble-like structures called synaptic vesicles. When this neuron is ready to send a signal to a second neuron, the membrane that encloses the synaptic vesicle fuses with the cell membrane that surrounds the neuron. This involves SNARE proteins in the vesicle membrane interacting with similar proteins in the cell membrane to form a SNARE complex, which then proceeds to 'zip' the two membranes together.

Other proteins are involved in the fusion process and the release of the neurotransmitters. For example, complexins bind to SNARE proteins during the formation of the SNARE complex in order to temporarily halt the fusion process. This 'clamping' interaction ensures that the neurotransmitters are released at the appropriate time.

Researchers have proposed two different models of the clamping interaction. In the *trans* clamping model a region in the complexins called the accessory helix extends forward and clamps SNARE proteins that are present on the two membranes. An alternative model explains clamping in terms of electrostatic interactions between the accessory helix and the two membranes. These interactions are repulsive because the accessory helix and the membranes are all negatively charged.

Now Krishnakumar, Li et al.—including some of the researchers who first proposed the *trans* clamping model—have used a variety of biochemical techniques to re-examine the clamping interaction. These experiments support the idea that the accessory helix binds to and clamps a SNARE protein, as suggested by the *trans* clamping model. The results of recent in vivo experiments on fruit flies have also provided support for the *trans* clamping model, although further work is need to compare the models in both in vitro and in vivo systems. DOI: 10.7554/eLife.04463.002

of CPX_{acc} which allows the *trans* clamping interaction, and the cell-cell fusion assay (*Hu et al., 2003*) to functionally test the zig-zag model for CPX clamping (*Krishnakumar et al., 2011*; *Kümmel et al., 2011*). Recently, Rizo, Rosenmund, and colleagues (*Trimbuch et al., 2014*) have re-examined the clamping interaction of CPX and have raised concerns regarding the interpretation of the ITC and FRET data and the use of the cell-cell fusion assay as an in vitro system to study CPX clamping (*Krishnakumar et al., 2011*; *Kümmel et al., 2011*). Here we address these concerns and argue that the *trans* clamping model we had previously proposed remains relevant.

Results

ITC experiments

In our earlier paper we used ITC experiments to confirm that the CPX_{acc} interacts with the t-SNARE in the truncated pre-fusion SNARE complex (SNARE∆60) as predicted by the X-ray crystal structure (Kümmel et al., 2011). To measure this interaction, we blocked the central helix binding site by prebinding the SNARE $\Delta 60$ complex with a truncated form of CPX (CPX-48; residues 48–134) before titration. In the recent report by Trimbuch et al. (2014) the authors suggest that the 1.5 molar excess of the CPX-48 that was used to block the CPX_{cen} binding does not saturate the central helix binding site and the heat observed upon addition of CPX to blocked SNAREΔ60 arises from the completion of CPX_{cen} binding rather than from interactions involving the CPX_{acc}. This was primarily based on their ITC data which showed that CPX-47 (CPX47–134) binds to truncated complex SNARE∆60 with an affinity constant (K_d) = 2.39 \pm 0.19 μ M and to non-truncated SNARE complex with K_d = 339 \pm 9 nM (Trimbuch et al., 2014). The binding constant for full-length CPX and a non-truncated SNARE complex is reported to be ~20 nM (Pabst et al., 2002) and, given that CPX-48 has an intact central helix including all SNARE-interacting residues (residues 48, 52, 69, and 70) (Chen et al., 2002), the expectation would be that CPX-48 and full-length CPX bind to the non-truncated SNARE complex with similar affinities. This discrepancy prompted us to repeat their ITC experiments and, under our experimental conditions, the K_d for CPX-48 binding to the post-fusion SNARE complex was 43 \pm 7 nM (Figure 1A, Table 1), much closer to the value reported for the full-length CPX (Pabst et al., 2002). We found further that CPX-48 bound the pre-fusion SNARE Δ 60 with a K_d = 457 ± 47 nM (*Figure 1B*, Table 1), or about five times more tightly than reported by Trimbuch et al. (2014). These binding



Figure 1. Characterization of interaction of complexin (CPX)-48 with pre- and post-fusion SNARE complex by isothermal titration calorimetry. Representative thermograms of CPX-48 titrated into post-fusion SNARE (**A**) or pre-fusion SNARE Δ 60 complex (**B**). The solid lines represent the best fit to the corresponding data points using non-linear least squares fit with the one-set-of-sites model. The results of the fit from 2–3 independent trials are shown in **Table 1**.

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constants ensure the near saturation (~96%) of the central helix binding site under the conditions (1.5 molar excess of CPX-48) reported in *Kümmel et al. (2011)*. Thus, with the blocked SNARE Δ 60, only the interaction between CPX_{acc} and t-SNARE would be measured. This is supported by the fact that the interaction affinity could be modulated by the mutations in CPX_{acc}. Hydrophobic mutations (D27L, E34F, R37A) increased the binding affinity (approximately eightfold stronger than the wild-type, *Table 1*), while the introduction of charged residues (A30E, A31E, L41E, A44E) abolished the interaction (*Kümmel et al., 2011*).

To restore confidence in our ITC data reported in *Kümmel et al. (2011)*, we also repeated the ITC binding experiments using 2.5–3-fold molar excess of CPX-48 to completely block the CPX_{cen} binding (\geq 99%). We found that CPX binds to the blocked SNARE Δ 60 with a binding affinity of 15.2 ± 1.4 μ M (*Figure 2A, Table 1*), matching well the K_d ~16 μ M reported in *Kümmel et al. (2011)*. We note that in these experiments as well as those reported in Kümmel et al. we titrated full-length CPX (residues 1–134) into the blocked SNARE Δ 60, and not the minimal functional domain (residues 26–83) as we had implied ("we used a complexin construct comprising both the central and accessory helices [residues 26–83]" [*Kümmel et al., 2011*]), and we apologize for this reporting error. We have now additionally carried out the ITC experiments with the minimal functional domain (CPX26–83) and find that this truncated version also binds to the blocked SNARE Δ 60, albeit with slightly weaker affinity (K_d = 23.9 ± 0.1 μ M) compared with full-length CPX (*Figure 2B, Table 1*). Taken together, the data strongly support our earlier conclusion that the ITC binding studies carried out with CPX titrated into

Table 1. Affinity constants (K_d) for complexin (CPX) binding to SNARE complexes measured by isothermal titration calorimetry

Complexin	Binding partner	Binding affinity (K _d)	Reference
CPX1-134	Ternary SNARE	19 nM	Pabst et al. (2002)
CPX48–134	Ternary SNARE	43 ± 7 nM	This study
CPX48–134	SNAREA60	457 ± 47 nM	This study
CPX1-134	Blocked SNARE∆60 (1.5-fold excess of CPX48–134)	16 μM	Kümmel et al. (2011)
Super-clamp CPX1–134 (D27L E34F R37A)	Blocked SNARE Δ 60 (1.5-fold excess of CPX48–134)	2 µM	Kümmel et al. (2011)
CPX1-134	Blocked SNARE Δ 60 (3-fold excess of CPX48–134)	15.2 ± 1.4 μM	Current study
CPX26-83	Blocked SNARE Δ 60 (3-fold excess of CPX48–134)	23.9 ± 0.1 μM	Current study
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blocked SNARE Δ 60 correctly reflect the binding of CPX_{acc} to t-SNARE. Consistent with this, we have recently also been able to characterize the binding of mammalian CPX_{acc} to *Drosophila* t-SNAREs using blocked *Drosophila* pre-fusion SNARE complex (*Cho et al., 2014*). Mutations in the CPX_{acc} predicted to enhance or decrease the binding of CPX_{acc} to t-SNARE exhibit corresponding binding profiles in ITC experiments (*Cho et al., 2014*) in support of the *trans* insertion model (*Kümmel et al., 2011*).

Further, to directly monitor the multiple binding modes of CPX to the pre-fusion SNAREpin, we carried out new ITC experiments where we titrated full-length CPX into unblocked SNARE Δ 60. As our initial ITC data had suggested that the accessory helix of super clamp CPX (residues 1–134 with D27L, E34F, R37A; scCPX) binds to SNARE Δ 60 with ~10× higher affinity (*Table 1*) than wild-type (*Kümmel et al., 2011*), we used scCPX for this analysis. As shown in *Figure 3A*, titration of scCPX into the unblocked SNARE Δ 60 results in a thermal graph characteristic of a reaction involving multiple binding sites, demonstrating that CPX has more than one binding site per SNARE Δ 60 complex. The data can be best approximated using the independent thermodynamic parameters for the CPX_{cen} and CPX_{acc} interaction, with the assumption that both the truncated SNARE Δ 60 and scCPX are bivalent (*Figure 3B*). We observed a qualitatively similar titration curve for wild-type CPX (*Figure 3C*) but, since the CPX_{acc} interaction with SNARE Δ 60 is much weaker, the fitting with multiple binding sites was not resolved in detail.

We suspect that experimental factors such as the buffer conditions, purification methodology (i.e., presence or absence of affinity tags), and the method used to determine the protein concentration (Bradford/BCA or A280) might contribute to the variability in the ITC data between the two papers. The latter is of particular significance since the quantitation of small proteins/peptide (<7 kDa) is highly dependent on the method used. However, we are unable to pinpoint the differences in protocol between the two studies since several relevant experimental details (e.g., whether affinity tags were present or what method was used for protein quantification) are not described in Trimbuch et al. Another discrepancy with regard to the ITC measure may derive from the way the blocked SNARE Δ 60 complex was assembled. In our experience, assembly of SNARE complex with VAMP60 or similarly truncated VAMP using only the concentration-dilution cycles as carried out in the Trimbuch et al. report results in heterogeneous samples, with non-productive aggregates and un-assembled components not very effectively removed by the concentration cycle. In our experimental regimen we always purified the truncated SNARE complex on a Superdex 75 gel filtration column to ensure good quality of the assembled complexes. Column-purified SNAREA60 complexes were subsequently incubated with gel-filtration purified CPX-48 to form the blocked SNARE $\Delta 60$ complex used in the ITC experiments. We also note that Pabst et al. (2002) used a more stringent purification protocol (Mono-Q purification) even though they used a different SNAP25 construct for the preparation of the SNARE complexes for their ITC studies and report K_d values similar to our findings (Pabst et al., 2002). The samples used by Trimbuch et al. for the NMR analysis appear to be homogeneous, however.



Figure 2. Interaction of complexin accessory helix (CPX_{acc}) with the t-SNARE groove for full-length and truncated (residue 26–83) CPX characterized by isothermal titration calorimetry. Full-length (**A**) or CPX26–83 (**B**) were titrated into pre-fusion SNARE Δ 60 complex with the CPX central helix (CPX_{cen}) binding site blocked with CPX-48 to exclusively measure the CPX_{acc}-t-SNARE clamping interaction. The solid lines represent the best fit to the corresponding data points using non-linear least squares fit with one-set-of-sites-model and results of the fit are shown in **Table 1**. All experiments were conducted in triplicate and a representative thermogram is shown. DOI: 10.7554/eLife.04463.005

FRET analysis

We used FRET analysis to establish that the angled conformation of CPX_{acc} also occurs in solution and is not dictated by crystal packing (Kümmel et al., 2011). We placed the donor probe on SNAP25, acceptor probe on CPX_{acc}, and used donor quenching to track the positioning of the CPX_{acc} in the pre- and post-fusion SNARE complex (Krishnakumar et al., 2011; Kümmel et al., 2011). The FRET data were consistent with the CPX_{acc} locating parallel to the SNARE complex in the fully-assembled SNARE complex, but moving away from the SNAREs in the pre-fusion half-zippered complex (Krishnakumar et al., 2011; Kümmel et al., 2011). In Trimbuch et al. (2014), based on NMR analysis, it was stated that the CPX_{acc} helix is poorly structured even when bound to the SNARE complex and exhibits higher flexibility with c-terminal truncation of VAMP. The authors therefore suggested that the low FRET state we observed in the pre-fusion SNAREpin and assign to the 'angled' conformation can be explained by the enhanced flexibility of the CPX_{acc} in this complex. Even though this is not in contradiction to the trans clamping model we have proposed, there are several lines of evidence (Krishnakumar et al., 2011; Kümmel et al., 2011) arguing against this interpretation. (1) We observe a well-defined 'high' and 'low' FRET state for CPX bound to the post- and pre-fusion SNARE using two different FRET pairs with different R₀ values (Stilbene/Bimane, R₀ ~27Å and Bimane/Oregon Green, $R_0 \sim 38A$), and the FRET distances in these CPX–SNARE complexes match very well with the predicted



Figure 3. Isothermal titration calorimetry indicates multivalent interactions between SNAREΔ60 and CPX. (**A**) Calorimetric titration of super-clamp complexin (scCPX; residues 1–134, with D27L, E34F, R37A mutations) into prefusion SNAREΔ60 complex describes a multi-site interaction of CPX. The solid lines represent the predicted binding thermogram assuming that both scCPX and truncated SNAREΔ60 are bivalent with well-defined independent thermodynamic parameters describing CPX central helix and CPX accessory helix binding (**B**). (**C**) Representative thermogram of full-length wild-type CPX (residues 1–134) titrated into unblocked SNAREΔ60. DOI: 10.7554/eLife.04463.006

distance in the pre- and post-fusion crystal structures (*Kümmel et al., 2011*). Further, a study by Choi et al. suggests that a completely unstructured CPX would result in a higher FRET signal than a more structured form (*Choi et al., 2011*), so the decrease in FRET efficiency for CPX bound to SNARE $\Delta 60$ is not consistent with increased disorder. (2) For both the FRET pairs tested, acceptor placed at residue 31 on CPX_{acc} shows weaker FRET compared with residue 38, consistent with the idea that the CPX_{acc} extends away from the pre-fusion SNARE complex, with dye on residue 31 locating further away from the donor (on SNAP25) than the dye on residue 38 (*Kümmel et al., 2011, Figure 4*).



Figure 4. Bimane–Oregon green fluorescence resonance energy transfer (FRET) experiments probing the effect of flexibility on complexin (CPX) orientation in pre- and post-fusion CPX–SNARE complexes. The FRET labeling positions were residue 193 on SNAP25 (Bimane) and residue 38 on CPX (Oregon green). Fluorescence emission spectra of Bimane only (black), Oregon green only (grey), and Bimane–Oregon green labeled CPX–SNARE complexes containing VAMP2 (residues 25–96, orange), VAMP60 (residues 25–60, blue) confirms the angled (low FRET) configuration of CPX accessory helix (CPX_{acc}) in the pre-fusion SNARE complex (VAMP60). The near complete loss of FRET for the flexible CPX mutant (helix breaking GPGP insertion between CPX_{acc} and CPC central helix (CPX_{cen}); CPX–GPGP, dashed lines) compared with the wild-type (WT, solid lines) in both pre- and post-fusion SNARE complexes shows that the difference in FRET signal observed with WT CPX is not due to increased flexibility of the CPX_{acc} in the pre-fusion complex. FRET distances in these CPX–SNARE complexes determined from the quenching of the donor (Bimane) fluorescence is shown (table) and standard deviations are reported from 2–3 independent experiments.

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(3) In the accompanying paper (*Krishnakumar et al., 2011*), we tested the effect of VAMP truncation on the orientation of CPX_{acc} and observed the same well-defined low FRET state (~15% FRET) for CPX bound to SNARE Δ 69, SNARE Δ 65, or SNARE Δ 60 complexes (*Krishnakumar et al., 2011, Figure 1B*). But NMR analysis for same or similarly truncated SNARE complexes showed equally dramatic enhancement of local flexibility in CPX_{acc} with increasing truncation on the VAMP c-terminus (SNARE Δ 68 versus SNARE Δ 62 versus SNARE Δ 60 in *Trimbuch et al. (2014), Figure 2C,F*). (4) Finally, the FRET signal we observed in the 'low' FRET state is higher than the FRET signal obtained when we used a CPX construct with enhanced flexibility (CPX–GPGP) in which a helix-breaking GPGP linker was inserted between the central and accessory helices of CPX for either SNARE Δ 60 or the post-fusion SNARE complex (*Kümmel et al., 2011, Figure 4D*). Trimbuch et al. argued that, since we observe only a small change in FRET signal for CPX–GPGP compared with CPX–SNARE Δ 60, the results obtained with the GPGP mutant are not conclusive. To address this concern we tested CPX–GPGP bound to SNARE Δ 60 and SNARE complex using the medium range Bimane/Oregon green FRET pair ($R_0 \sim 38$ Å). As shown in *Figure 4*, the GPGP insertion results in a dramatic decrease in the FRET signal, with the CPX_{acc} locating further away from both SNARE Δ 60 and the SNARE complex. These data suggest that the increased flexibility does not explain the decreased FRET signal in the pre-fusion truncated SNARE complex.

Thus, taken together, our data are consistent with our earlier conclusion that the low FRET state corresponds to a defined angled configuration on the CPX_{acc} in the pre-fusion complex. This is corroborated by functional data showing that the inhibitory function of CPX requires a fully-folded and rigid CPX helix (*Radoff et al., 2014*), and introducing flexibility within the minimal functional domain of CPX (residue 26–83) results in a reduction or loss of clamping ability corresponding to the degree of instability introduced (*Kümmel et al., 2011; Cho et al., 2014; Radoff et al., 2014*).

NMR analysis

Using NMR, Trimbuch et al. were unable to detect any interaction between a peptide corresponding to CPX_{acc} only (CPX26–48) and SNARE Δ 60 (*Trimbuch et al., 2014*). This experiment is in agreement with our ITC measurements, which did not detect an interaction between SNARE Δ 60 and CPX26–48 alone (unpublished). We suspect that the conformations sampled by CPX26–48 are dependent on its protein context, and this might account for the different binding affinities observed for the shorter (CPX26–48) compared with the longer (CPX26–83 and CPX1–134) constructs. For example, our circular dichroism (CD) measurements showed that CPX_{acc} (residues 26–48) by itself is unfolded, but CPX26–83 forms a stable α -helical structure (*Figure 5A*).

However, Trimbuch et al. (2014) also did not observe any dramatic shifts and/or broadening of the CPX_{acc} cross-peaks, as would be expected for insertion into the t-SNARE groove (Kümmel et al., 2011), when they used CPX26-83. We note that the prediction from our model is that the trans CPX_{acc}-t-SNARE interactions would lead to the formation of CPX/SNARE∆60 polymers, which would not be visible in NMR studies due to line broadening. Thus, if there is a trans interaction, then the NMR technique may not be well suited for its interrogation. This cross-linked high-order oligomeric state is evident qualitatively in the ITC experiments where we titrated full-length CPX into unblocked SNARE $\Delta 60$ (Figure 3). Dynamic light scattering (DLS) analysis (Figure 5B, black circles) similarly suggests that CPX26-83/SNAREA60 complexes begin to oligomerize at the concentration regime used in Trimbuch et al. (25–50 μ M). The DLS data are consistent with a model predicting the size of the oligomers (see 'Materials and methods' for details on the model, Figure 5B, black line) which incorporates the K_d of \sim 25 μ M obtained from our ITC analysis of the CPX26-83 and blocked SNARE $\Delta 60$ interaction. In contrast, we observe only a small change in the apparent size if we disrupt the binding of CPX_{acc} to t-SNARE either by introducing non-clamping mutations (ncCPX26-83; A30E, A31E, L41E, A44E) in CPX_{acc} (Figure 5B, blue circles) or by blocking the t-SNARE groove using a fully assembled ternary SNARE complex (Figure 5B, green circles). Based on our oligomerization model, this small change in the particle size could only be fitted by assuming a very low affinity interaction $(K_d \ge 250 \ \mu M)$ such as might arise from non-specific aggregation (*Figure 5B*, blue and green lines).

We suspect that the concentrations used in the NMR studies (~25–50 μ M) (*Trimbuch et al., 2014*), which correspond to the lower end of the range of concentration used for NMR studies, were adjusted to avoid oligomerization and line broadening effects. Because small proteins are difficult to quantitate and because our K_d values, which depend on concentration measurements for their accuracy, are only approximate, it is plausible that, under the NMR conditions of Trimbuch et al., most of the CPX–SNARE Δ 60 complexes were monomeric, with only a small fraction in an oligomeric form. Thus, under the experimental conditions used by Trimbuch et al., CPX_{acc} might not have an appreciable interaction with the t-SNARE groove, and the minor bound fraction would not be visible in NMR spectra owing to the large size of the cross-linked oligomers, assuming that there is no exchange between the monomeric and oligomeric forms of CPX–SNARE Δ 60. Therefore, we believe that the negative NMR data in Trimbuch et al. do not preclude a *trans* interaction between CPX and the SNARE Δ 60 complex.



Figure 5. Circular dichroism (CD) and dynamic light scattering (DLS) experiments. (**A**) CD spectra of the complexin (CPX) accessory domain (CPX26–48) and the CPX minimal functional domain (CPX26–83). The continuous CPX construct (26–83, blue) with uninterrupted accessory and central domain shows the characteristics of an α -helical protein. In contrast, the isolated CPX accessory domain (26–48) shows little secondary structure and appears mostly unfolded. This may explain why no interaction between the CPX accessory region and SNARE Δ 60 was observed by NMR. (**B**) DLS analysis showing the formation of the high-order oligomers of CPX–SNARE Δ 60 in the concentration Figure 5. continued on next page

Figure 5. Continued

range used in the NMR experiments in **Trimbuch et al. (2014)**. Experimental average particle radius of pre-formed CPX26–83/SNAREΔ60 (black dots), ncCPX26–83/SNAREΔ60 (blue dots), and CPX26–83/SNARE (green dots) at varying concentration is shown. The solid lines (same color scheme) represent the average gyration radius of the oligomers calculated from the semi-quantitative model described in 'Materials and methods'. DOI: 10.7554/eLife.04463.008

In vivo analysis

In Trimbuch et al., the authors tested the CPX_{acc}/t -SNARE insertion model using the autaptic neuronal culture derived from CPX triple KO mice (CPX TKO). They found that the super clamp and non-clamp CPX mutations, which are predicted to enhance or decrease CPX_{acc} -t-SNARE binding, respectively, did not have the expected effect on evoked or spontaneous release. Based on this, they concluded that functional data do not support the CPX_{acc} insertion/zig-zag model (*Trimbuch et al., 2014*) and raised concerns regarding the use of the cell-cell fusion assay (*Giraudo et al., 2006, 2009*; *Krishnakumar et al., 2011*; *Kümmel et al., 2011*) as an in vitro system to study CPX clamping.

These results and conclusions must be viewed with caution since the studies in knockout neurons, particularly in autaptic neuronal cultures, reveal a more restricted role of CPX in the neurotransmitter release (Reim et al., 2001; Xue et al., 2007, 2008; Yang et al., 2013). Specifically, the inhibitory function of CPX has not been observed in the autaptic system: that is, both evoked and spontaneous release are reduced in CPX TKO mice (Reim et al., 2001; Xue et al., 2008; Trimbuch et al., 2014). In contrast, other preparations like rodent mass cultured neurons with reduced CPX expression and invertebrates lacking CPX exhibit reduced evoked release and enhanced spontaneous release (Huntwork and Littleton, 2007; Yang et al., 2010; Cho et al., 2014). Therefore, these systems may be more relevant to examine the mechanism of CPX function in regulating vesicle fusion, particularly in its ability to inhibit and regulate spontaneous release. In rescue experiments with rodent mass cultured neurons in which CPX was knocked down, CPX mutants that either enhanced or disrupted the CPX_{acc}-SNARE interactions (super-clamp and poor clamp, respectively) resulted in corresponding reductions and enhancements in spontaneous release compared with wild-type CPX controls (Yang et al., 2010). Recent experiments at the Drosophila neuromuscular junction, which in the absence of CPX exhibits large increases in spontaneous release, directly examined the role of a CPX zig-zag array in regulating spontaneous release. Using genetic rescue approaches, CPX mutants predicted to prevent the formation of the zig-zag array (CPX-GPGP) disrupt the ability of CPX to clamp spontaneous release, while mutations predicted to enhance the CPX_{acc}-t-SNARE interaction (super-clamp mutation) exhibit a strong clamping ability (Cho et al., 2014). These results taken together are consistent with a CPX cross-linking model mediating the CPX clamping function to regulate spontaneous release, even though recent reports suggest that the inhibitory and activating functions of CPX may be separable (Yang et al., 2010; Cho et al., 2014; Lai et al., 2014). In addition, these in vivo studies (Yang et al., 2010; Cho et al., 2014) support many of the observations made first with the cell-cell fusion assay (Giraudo et al., 2006, 2008, 2009; Krishnakumar et al., 2011; Kümmel et al., 2011; Li et al., 2011), underscoring the validity and the relevancy of the in vitro cell-cell fusion assay to study CPX clamping.

Electrostatic hindrance model

As an alternative to the insertion/zig-zag model, Trimbuch et al. advanced an 'electrostatic hindrance model' for CPX clamping (*Trimbuch et al., 2014*). This was based on the finding that increasing or decreasing the net negative charge on CPX_{acc} inhibits or stimulates neurotransmitter release, respectively (*Trimbuch et al., 2014*). We note that a number of both in vitro and in vivo studies on CPX_{acc} modification argue strongly against the electrostatic hindrance model (*Yang et al., 2010*; *Kümmel et al., 2011*; *Cho et al., 2014*; *Radoff et al., 2014*). Introducing two (L41E, A44E) or four (A30E, A31E, L41E, A44E) negatively charged mutations in the mammalian CPX_{acc} has been shown to entirely abolish or severely diminish the interaction of CPX_{acc} with the mammalian or *Drosophila* t-SNARE, respectively (*Kümmel et al., 2011*; *Cho et al., 2014*). Consistent with this finding, the similar mutations have been shown to abrogate the clamping function of CPX in both in vitro cell–cell fusion assay (*Kümmel et al., 2011*) and in vivo rescue experiments with knockdown neurons (*Yang et al., 2010*). Further, in contrast to the electrostatic model, increasing the hydrophobicity of CPX_{acc} has been shown to enhance the clamping function of CPX (*Yang et al., 2011*; *Kümmel et al., 2011*; *Cho et al., 2010*; *Kümmel et al., 2010*; *Kümmel et al., 2011*; *Cho et al., 2010*; *Kümmel et al., 2011*; *Cho et al., 2010*; *Kümmel et al., 2011*; *Cho et al., 2010*; *Kümmel et al., 2010*; *Kümmel et al., 2011*; *Cho et al., 2010*; *Kümmel et al., 2011*; *Cho et al., 2014*).

Discussion

We believe that the new in vitro data presented above and physiological data published recently (Cho et al., 2014) should dispel the concerns raised by Trimbuch et al. regarding the plausibility of the insertion/zig-zag model. Specifically, ITC binding experiments with blocked SNARE complexes, where a 2.5–3-fold excess of blocking peptide was present, have corroborated the interaction of CPX_{acc} with the t-SNARE groove in a pre-fusion (truncated) SNARE complex as observed in the crystal structure (Figure 2 and Kümmel et al., 2011). Additionally, titration of scCPX into unblocked truncated SNARE complex revealed multiple binding sites consistent with independent binding sites for CPX_{cen} and scCPX_{acc}, thus describing the essential feature of the trans clamping interaction of CPX (Figure 3). FRET analysis (using two FRET pairs with different R₀ values) with 'flexible' CPX construct (CPX-GPGP) has clearly demonstrated that the low FRET state corresponding to the angled conformation, which describes the trans clamping interaction, is due to discrete conformational change and not because of increased CPX flexibility (Figure 4 and Kümmel et al., 2011). Lastly, the physiological relevance of the insertion model was established by independent genetic rescue experiments in the Drosophila neuromuscular junction, wherein mutations in CPX that are predicted to prevent the formation of the SNAREpin array were found to disrupt the ability of CPX to clamp spontaneous release (Cho et al., 2014).

All of our data are consistent with a model in which CPX_{acc} from one SNAREpin interacts in trans with the t-SNARE groove of a second SNAREpin. As we emphasized in Kümmel et al., we do not propose that the interactions are necessarily exactly as observed in a single crystal structure for which scCPX was used and where hydrophobic accessory helix residues (which are hydrophilic in wild-type CPX) interact with hydrophobic residues in a partially assembled SNARE complex. Details of the interaction between wild-type CPX_{acc} and t-SNARE must be different (they are also weaker, explaining why the wild-type CPX clamps less well), and we can well imagine a scenario in which wild-type CPX_{acc} interacts with a slightly different surface of the assembling SNARE complex. In fact, we found that a single mutation in the CPX_{acc} (F34M) results in two distinct binding interfaces for scCPX, but both giving rise to the same zig-zag topology. Both interactions involved the same face of CPX_{acc} and t-SNARE, although the binding site on t-SNARE was extended by two helical turns for the mutant (Kümmel et al., 2011). The consideration that interactions of wild-type and scCPX and the SNARE complex are not identical does not, however, exclude the possibility that a trans interaction of some sort is responsible for clamping. We also suspect that yet unknown additional interactions in the prefusion complex, which are not represented in our crystal structure, may further stabilize the trans clamping interaction (Cho et al., 2014; Radoff et al., 2014). We do not consider the trans clamping model proven, but it remains the most well-defined model for which a good amount of evidence is available.

Materials and methods

Plasmid constructs and protein purification

The constructs used in this study are GST-PreScission-VAMP2Δ60 (human VAMP2 residues 29–60), pET15b-oligohistidine-thrombin-VAMP2 (human VAMP2 residues 29–96), pET28-oligohistidine-synataxin1A (containing rat syntaxin 1a residues 191–265), oligohistidine-SUMO-SNAP25N (containing human SNAP25A residues 7–82), oligohistidine-SUMO-SNAP25C (containing human SNAP25A residues 141–203) and pET15b-oligohistidine-thrombin-CPX (containing human complexin1 residues 1–134); residues 1–134 with flexible GPGP insert between residues 49–50 (CPX–GPGP), residues 26–83 (CPX26–83); residues 26–83 with non-clamping mutations A30E, A31E, L41E, A44E (ncCPX26–83). Truncated CPX constructs, including residues 48–134 (CPX-48), residues 26–48 (CP26–48) was purified as oligohistidine-SUMO constructs. All constructs were expressed and purified as described previously (*Krishnakumar et al., 2011; Kümmel et al., 2011*). To ensure high quality, all proteins were purified on a High-load Superdex 75 (16/60, GE Healthcare; Piscataway, NJ) gel filtration column.

ITC analysis

ITC experiments were carried out as described previously (*Krishnakumar et al., 2011*; *Kümmel et al., 2011*; *Cho et al., 2014*). To assemble the post-fusion and pre-fusion SNAREΔ60 complex, Syntaxin 1a, SNAP25N, SNAP25C, and VAMP2 or VAMPΔ60 were mixed (molar ratio of 1:1.2:1.2:1.2 for the post-fusion complex and 1:1.2:1.2:1.6 for pre-fusion SNAREΔ60 complex) and incubated overnight at

4°C. The assembled complexes were purified from non-productive aggregates and unassembled components by gel filtration (High-load Superdex 75 16/60, GE Healthcare). To form the blocked SNAREA60 complex, purified SNAREA60 complex was mixed with 2.5 molar excess of CPX-48 and incubated overnight at 4°C to ensure complete binding. Then, we titrated CPX-48 back into this initial mixture of SNARE $\Delta 60$ and CPX48-134. If there was heat signal, we added more CPX-48 to the mixture. This process was repeated until negligible residual heat signal was observed (control experiment). Then we titrated full length CPX1-134 into the exact same mixture of blocked SNARE∆60 as used in the control experiment. By comparing the full length CPX titration with the control titration, we determined the binding interaction between CPX N-terminus and C-terminal unzippered t-SNARE. To ensure buffer uniformity to measure the weak interactions, CPX variants and the different SNARE complexes were extensively dialyzed (4L for 4hr followed by another 4L overnight) into the same phosphate buffered saline (PBS) buffer (pH 7.4, 137 mM NaCl, 3 mM KCl, 10 mM sodium phosphate dibasic, 2 mM potassium phosphate monobasic, 0.25 mM TCEP) before the ITC analysis. The concentrations of dialyzed proteins were determined by bicinchoninic acid (BCA) protein assay kit (Thermo Scientific; Waltham, Ma) and/or Bradford assay (Bio Rad; Hercules, CA) with bovine serum albumin (BSA) as the standard. ITC experiments were performed with a Microcal ITC200 (Malvern Instruments, UK). Typically, ~200 µl of SNARE solution was loaded into the sample cell and \sim 60 μ l of CPX solution was loaded into the syringe. The protein concentrations used for the titration were as follows: 110 µM CPX-48 titrated into 5.8 µM SNARE (Figure 1A); 210 μM CPX-48 titrated into 14 μM SNAREΔ60; ~360 μM CPX1-134 or CPX26-83 titrated into ~20 μM blocked SNAREΔ60 (Figure 2); 150 μM wild-type or scCPX1-134 titrated into 7.5 μM SNAREΔ60 (Figure 3). The heat change from each injection was integrated and then normalized by the moles of CPX in the injection. The thermographs were analyzed by non-linear least squares fit with the one-set-of-sites-model in Microcal Origin ITC200 software package to obtain the stoichiometric number (N), the molar binding enthalpy (ΔH), and the association constant (K_a). The equilibrium dissociation constant (K_d), the binding free energy (ΔG), and the binding entropy (ΔS) were calculated using the thermodynamic equations:

$$K_d = \frac{1}{K_a}.$$

$$\Delta G = \Delta H - T\Delta S = -RT\ln(K_a).$$

FRET analysis

FRET measurements were carried out as described previously (*Krishnakumar et al., 2011*; *Kümmel et al., 2011*). SNAP25 D193C was labeled with the donor probe, Bimane (Monochlorobimane, Invitrogen) and the scCPX Q38C was labeled with acceptor Oregon green (Oregon green 488 maleimide, Invitrogen). The proteins were labeled using 10x molar excess of dye in 50 mM Tris buffer, pH 7.4, containing 150 mM NaCl, 10% glycerol, and 1 mM TCEP. Following overnight incubation at 4°C, the excess dye was separated from the labeled proteins using a NAP desalting column (GE Healthcare). The labeling efficiency was calculated using $\varepsilon_{396} = 5300 \text{ L M}^{-1} \text{ cm}^{-1}$ for Bimane and $\varepsilon_{496} = 76,000 \text{ L M}^{-1} \text{ cm}^{-1}$ for Oregon green, and the protein concentration was measured using the Bradford assay with BSA as standard. Typically, the labeling efficiency was >95% for both Bimane-SNAP25 and Oregon green-CPX. The double-labeled CPX–SNARE complexes were assembled overnight at 4°C and purified by gel filtration on a Superdex 75 (10/30, GE Healthcare) gel filtration column. All fluorescence data were obtained on a Perkin-Elmer (Waltham, MA) LS55 luminescence spectrometer operating at 25°C. Excitation and emission slits of 5 nm were used in all measurements. Fluorescence emission spectra were measured over the range of 410–600 nm with the excitation wavelength set at 396 nm. The donor probe concentration was adjusted to 2 μ M in all samples.

Circular dichroism (CD) analysis

CD spectra of peptides corresponding to the CPX accessory domain alone (residues 26–48) or the minimal functional domain (residues 26–83) were recorded in PBS using Chirascan CD spectrometer (Applied Photophysics, UK) at 25°C from the range of 260 nm–198 nm at 1 nm bandwidth.

Dynamic light scattering (DLS) analysis

CPX-SNARE∆60 complexes containing wild-type CPX1-134 were assembled and purified as previously described (*Krishnakumar et al., 2011*; *Kümmel et al., 2011*). DLS experiments were

carried out on a DynaPro NanoStar instrument (Wyatt Technology; Santa Barbara, CA) at a wavelength of 663.76 nm operating at 4°C. Protein samples were centrifuged (10 min at 13,000×g) and data were collected using DynaPro disposable cuvettes. Autocorrelations for 20 s were collected over 15 acquisitions. Points were eliminated if the intensity fluctuated by more than 15% from the average. Data were analyzed with DYNAMICS 7.1.7.16 software (Wyatt Technology).

In DLS there is oligomerization of CPX–SNARE Δ 60. We can try to model this process as an equilibrium between bound and unbound CPX–SNARE Δ 60, wherein the bound state is generated by the interaction of CPX_{acc} with the t-SNARE groove. Thus, this equilibrium can be written as:

$$CPX - SNARE\Delta 60 + CPX - SNARE\Delta 60 \stackrel{N_d}{\leftrightarrow} CPX - SNARE\Delta 60 - CPX - SNARE\Delta 60.$$
(1)

which, in terms of concentration, gives:

$$[un - bound CPX - SNARE\Delta 60]^2 = Kd * [bound CPX - SNARE\Delta 60].$$
(2)

Keeping in mind that

$$[un - bound CPX - SNARE\Delta 60] + [bound CPX - SNARE\Delta 60] = [CPX - SNARE\Delta 60]_{initial},$$
 (3)

where $[CPX-SNARE\Delta 60]_{initial}$ is the initial concentration, the equilibrium concentrations can be calculated from (2) and (3) as:

un - bound CPX - SNARE
$$\Delta 60$$
] = $\frac{K_d \left(\sqrt{1 + \frac{4[CPX - SNARE\Delta 60]_{initial}}{K_d}} - 1\right)}{2}$. (4)

$$[\text{bound CPX} - \text{SNARE}\Delta 60] = [CPX - SNARE\Delta 60]_{initial} - \frac{K_d \left(\sqrt{1 + \frac{4[CPX - SNARE\Delta 60]_{initial}}{K_d}} - 1\right)}{2}.$$
 (5)

Then, the probability that an accessory helix is bound is given by:

$$p = \frac{[bound CPX - SNARE\Delta 60]}{[CPX - SNARE\Delta 60]_{initial}} = 1 - \frac{K_d \left(\sqrt{1 + \frac{4[CPX - SNARE\Delta 60]_{initial}}{K_d}} - 1\right)}{2[CPX - SNARE\Delta 60]_{initial}},$$
(6)

and the average number of monomers is equal to:

$$N = \frac{p}{1-p} + 1 = \frac{2[CPX - SNARE\Delta 60]_{init}}{K_d \left(\sqrt{1 + \frac{4[CPX - SNARE\Delta 60]_{init}}{K_d}} - 1\right)}.$$
(7)

We will model the gyration radius of an N-mer as:

$$= N^{0.5} r_0,$$
 (8)

where r_0 is the distance between two particles in the array and the reflection of monomer size. The r value is quantitative for long ideal polymers but is only semi-quantitative for non-ideal oligomers under consideration here.

The experimental results can be well approximated by **Equation 8** using $r_0 = 3.5$ nm and the CPX_{acc}-t-SNARE interaction described by $K_d = 25 \,\mu$ M for CPX26-83 titrated into blocked SNARE Δ 60. This r_0 value is reasonable considering the dimensions of the CPX-SNARE Δ 60 unit within the zig-zag array. Hence, the oligomerization we observe by DLS is consistent with the K_d values we measured by ITC (*Figure 2*). In support of the idea that the CPX_{acc}-t-SNARE clamping interaction results in the oligomeric state, CPX-SNARE complexes containing either ncCPX26-83 (A30E, A31E, L41E, A44E) or the full ternary SNARE complex show only a small change in the average particle radius. This behavior can be modeled using *Equation 8* with $r_0 = 3.8$ nm and only if we assume a very low affinity interaction, namely $K_d = 250 \,\mu$ M for ncCPX into SNARE Δ 60 and $K_d = 300 \,\mu$ M for CPX into the full ternary SNARE complex. This suggests that this change in particle size might be a result of non-

specific aggregation. Note that r_0 values for specific and non-specific oligomerization are not necessarily expected to be identical.

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Author ORCIDs

Daniel Kümmel, Daniel Küm

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