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#### Localization of the Sodium Dependent Cation-Chloride Cotransporter in Aedes aegypti

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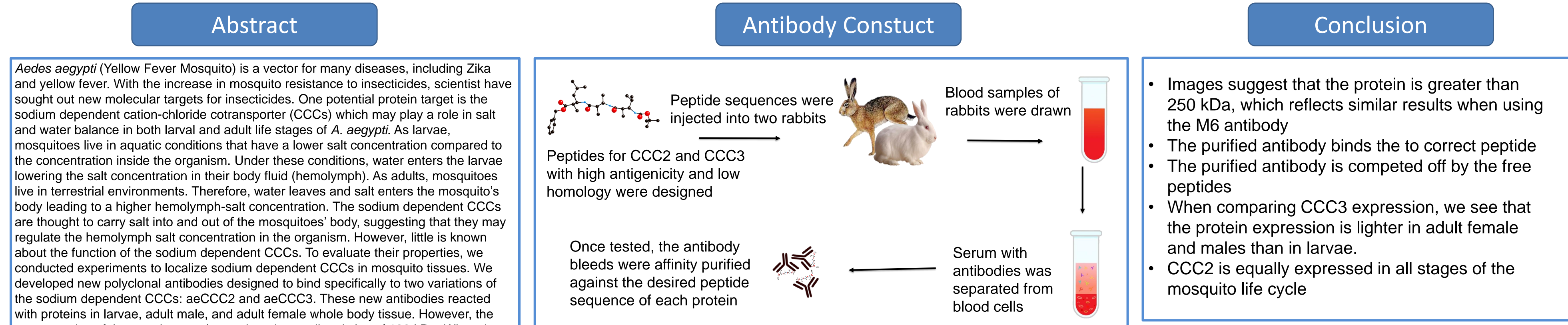
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# Localization of the Sodium Dependent Cation-Chloride Cotransporter in Aedes aegypti Kenyatta Viel '17 and Chris M. Gillen, Ph.D. Kenyon College Summer Science 2016



apparent size of the proteins was larger than the predicted size of 130 kDa. When the samples were exposed to urea, which disaggregates protein complexes, and PNGase F, which cleaves sugar residues off proteins, the apparent size of the protein detected by the antibodies did not decrease. We are now using these antibodies to localize aeCCC2 and aeCCC3 in larval and adult renal tissue, which are known to be involved in salt transport.

## Background

Mosquitoes have three versions of the Na<sup>+</sup> dependent cation-chloride cotransporter (aeCCC), which move ions across epithelial tissue in renal organs. However, limited research has been done on the function of each transporter and how it relates to the molecular physiology in this insect. Previous research conducted in the lab showed that the aeCCC1 is more phylogenetically related to the functionally characterized Na<sup>+</sup>, K<sup>+</sup>, 2Cl<sup>-</sup> cotransporters (NKCC). One study, has shown that the NKCC protein in *Drosophila* melanogaster is required for normal renal tubule function (Rodan et al. 2012). The renal tubule, like the Malpighian tubule, is used to secrete ions and water from the body and into the environment. Because these two proteins are closely related, aeCCC1 is probably found on the basolateral side of secretory tissue as seen in the Drosophila's NKCC. The aeCCC2 and the aeCCC3 are closely related to transporters in Anopheles gambiae, another mosquito species; however, very little research has been done on these transporters in mosquitoes or in other insects (Figure 1). Previous lab work has shown that aeCCC2 and aeCCC3 are more localized in the adult hindgut and in the larval anal papillae, respectively, which absorb ions and water from the environment to the body. This suggests that aeCCC2 and aeCCC3 may be absorptive cotransporters, found on the apical side of the hindgut and anal papillae. However, sequence similarity does not necessarily dictate physiology. In order to determine the functions of the CCC proteins, it is necessary to determine where each protein is more heavily expressed and what kind of environmental factors increase expression.

Figure 2. Flowchart of the process of creating polyclonal antibodies in rabbits

IEYSSLTMLQGV 956 INYVSLIMV-TL 920

---ALLYRGPG 933

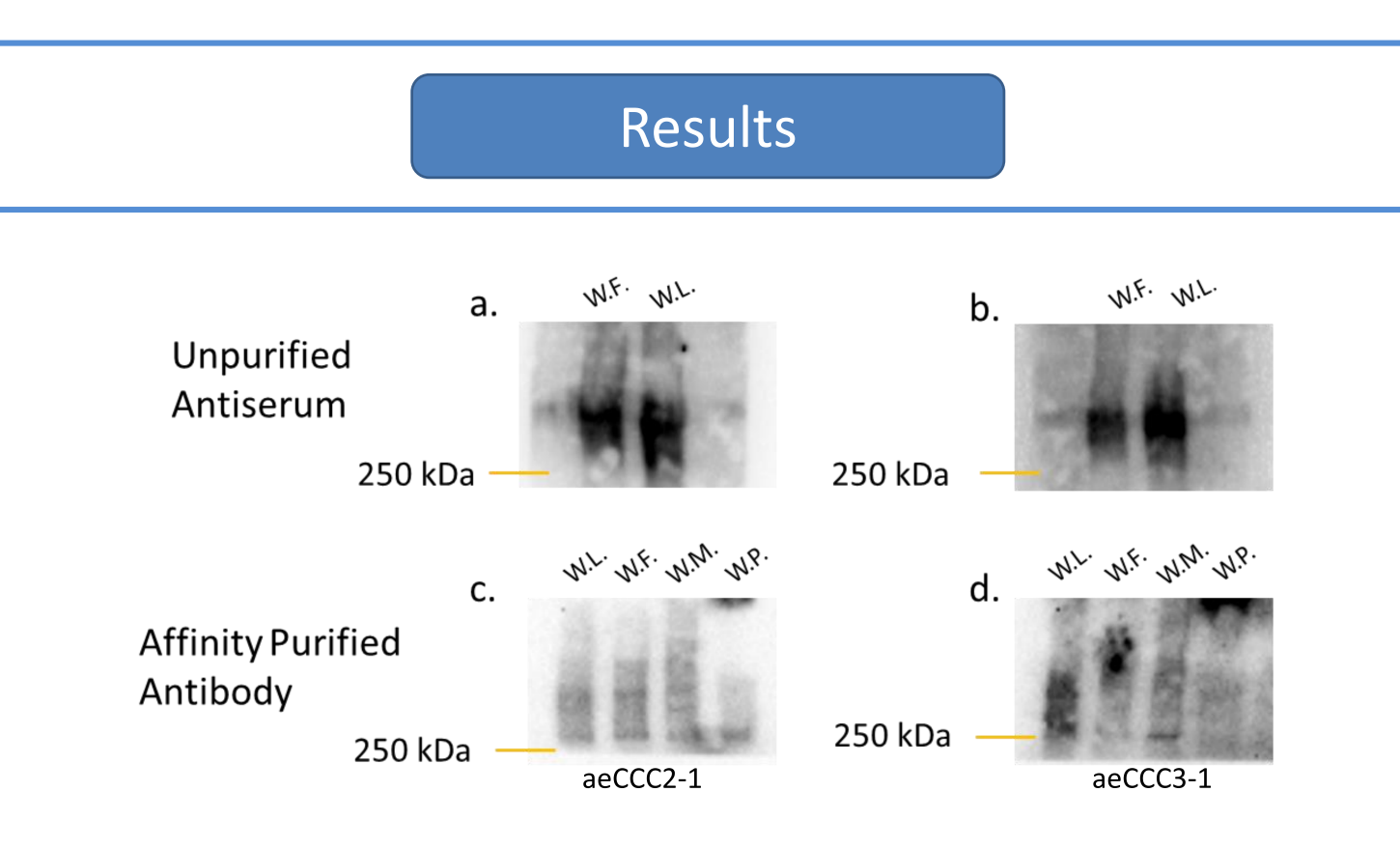
YKTKTYRQIR 1151

GNQTSVLTFY 1040 GNHQSVLTFY 1211

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msBSC	SYKVRSARARSDDEWLRGRKVRAFCSRVHGFSFEPGARALVQGSGVGRLAPNVLLMGY	726	msBSC	GLTILLPYIISQRSAWANCKLRIFALANRLHEMELEERNMANLLAKFRI
aeCCC2	SHRKRMEMNSEC <mark>NKFLEIRKIRGFYQPI</mark> DGLSFEEGVHALIQTSGVGKLSPNIVLMGY	737 2-2	aeCCC2	GLTMLIPYIISMRSKWARCKIRVFALTNRQLELEVEERNMANLLMKLRI
aeCCC3	NYKERKAYIQSGKKVLKDLKIKAFYSVLDGLPFDESVRAMIQSTGFGRLSPNILMVGY	719	aeCCC3	GLTILLPYIISTRSKWSECQIRVFALATQQTNVEEERENMTILLEKLRI
aeCCC1	SQKFRNYLQRKATDWFRRHKVKGFYTYVDDNEFETGARAAMQASGIGKLRPNLLLLGY	772	aeCCC1	GMPDPMDINA-KLLNETNNRSLKRTHNDP
hNKCC1	RQAMKEMSIDQAKYQRWLIKNKMKAFYAPVHADDLREGAQYLMQAAGLGRMKPNTLVLGF	888	hNKCC1	GLTLLIPYLLTTKKKWKDCKIRVFIG-GKINRIDHDRRAMATLLSKFRI
	: *::.* : : :* :*: ** :::*:			*: * : ::
msBSC	KSDWTTCPANDLVSYFNVLHTAFENRLAVAIVRVSGGLDYSAVVSEGAEEGAAGS	781	msBSC	TDPPQPETKALFDETIKKFTEESASPDCRISDMELQ
aeCCC2	KSDWMTCPVKDLLTYYNVLHDSFDCRMALAILRLPNGLDFSDLTTEIVTRTVPIPSTLKD	797	aeCCC2	TDAPRQET <mark>VDMHSKLLQHFTDNDGTQIP</mark> ISEHERM
aeCCC3	KQDWRTCGNAELHSYYNILHNAFDNRLALTILRLPNGLDCSHLVPPKPALDLADSALN	777	aeCCC3	SDKPQEAT <mark>IQSHKALLGTLVEGQETDVF</mark> VSASEQA
aeCCC1	KNDWRKCDSVELEQYFNVVHKALDMYLSVAILRVAKGLDYSQVLGEDTAAKQI	825	aeCCC1	GAELPKEVLDELTQFTSKRARVIDVYWLYISDAELL
hNKCC1	KKDWLQADMRDVD <mark>MYINLFHDAFDIQYGVVVIRLKEGLDISHLQGQEELLSSQ</mark>	941	hNKCC1	NTKPKKENIIAFEEIIEPYRLHEDDKEQDIADKMKEDEPW-RITDNELE
	*.**: * ** .:: .:*: *** * :			: *
msBSC	LTATSSSGELRVRRDGLIMHADSDLDIRDTQPKHNLSNLLTLTTSRSFTISECKEK	837	msBSC	LRELLLANSKDARLVVMSLPMPRKGSISAPLYMAWLEMMSRDLPPMLFV
aeCCC2	RSDNAISTISSDGMRTPRNLMHVDSNLNLDSLGGSANQLPGTPQ	841	aeCCC2	LREMLLEHSNEANLIVMSMPMPRLGTVSAPLYMSWLEMLTKGMPPFLLV
aeCCC3	IMSSMAYINTPQGLRIQKGLMPVNSNLDLHGMAPSSEQ <mark>ISVNVP</mark>	821 3-1	aeCCC3	LRELLQQYSKNASLIVLSMPIPRKGIVSAPLYMSWLEMLTKDMPPFLLV
aeCCC1	AETPRTLLHNDSSNDLAGQNKI-NSLHGSCDSLSRNISQGPEH	867	aeCCC1	LREYLLEHSKKSDLVVMTLPMPRKGVVSAPLYMAWLETLSQGLPPFLFV
hNKCC1	LDTSKPLSEKPIT	974	hNKCC1	LNELLKEHSSTANIIVMSLPVARKGAVSSALYMAWLEALSKDLPPILLV
	. * :			*.* * *. : ::*::*: * * :*: ***:*** ::: :**:*:*
msBSC	DKKKKERKPNDMHRQIVYNTASG <mark>LELSKFQLAQMSLFQKKQESGTLDVWWLYDDG</mark>	892	msBSC	S 1060
aeCCC2	MAPSPTVETSDIADDIIYSTRGGSCVPKEILDRIGVFQRKQPKGTIDVWWLYDDG	896	aeCCC2	S 1063
aeCCC3	NEAQDNKTTPEAAAPSQELNIFREKQPAGYIDVWWLYDDG	861	aeCCC3	S 1027
aeCCC1	NN-VHEKNEKNVS	897	aeCCC1	s 1041 M6 T4 aeCCC2
hNKCC1	HK-VEEEDGKTATQPLLKKESKGPIVPLNVADQKLLEASTQFQKKQGKNTIDVWWLFDDG	1033	hNKCC1	s 1212

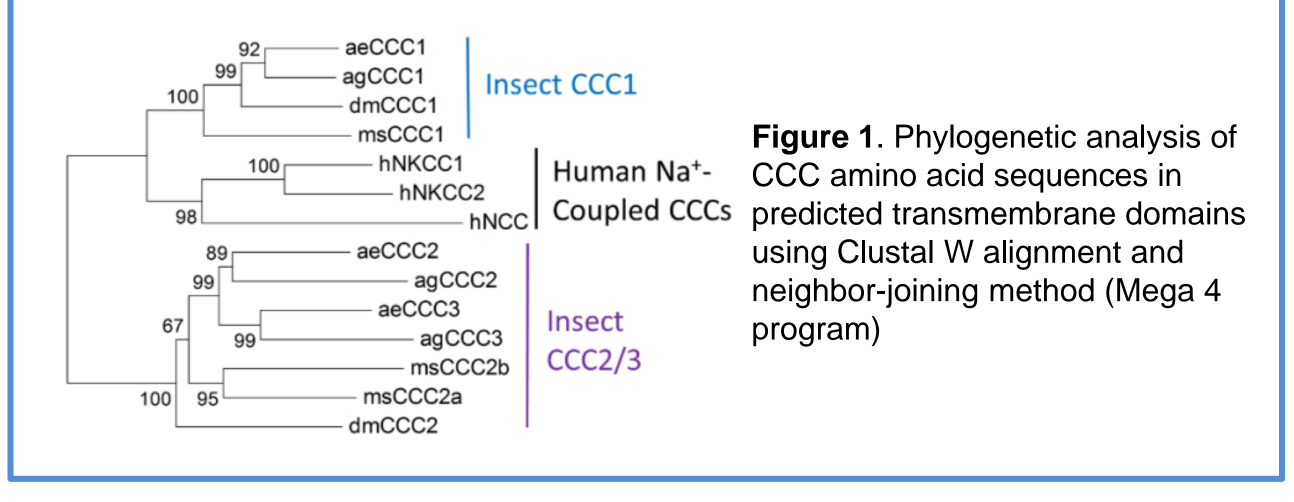
**Figure 3.** Amino acid sequence of antibodies M6 and T4 in comparison to the developed antibodies



Future Research

- Because these are initial results, more blots need to be produced in order to confirm protein expression between larvae and female adults
- The apparent size of the protein is larger than the predicted size of 130 kDa.
- Where are these proteins expressed in different tissues? We predict that CCC2 will be present in hindguts of larvae and adult female and that CCC3 will be present in anal papillae of larvae
- A peptide competition experiment was conducted with the bleeds. The figure suggest that the lower band was bound by the CCC2 antibodies. However, this experiment needs to reexamined with the affinity purified antibodies.

No peptides	Peptides for CCC2	Peptides for CCC3
W.F. W.L.	N.E. N.F.	N.F. N.L.
- 250	-	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2

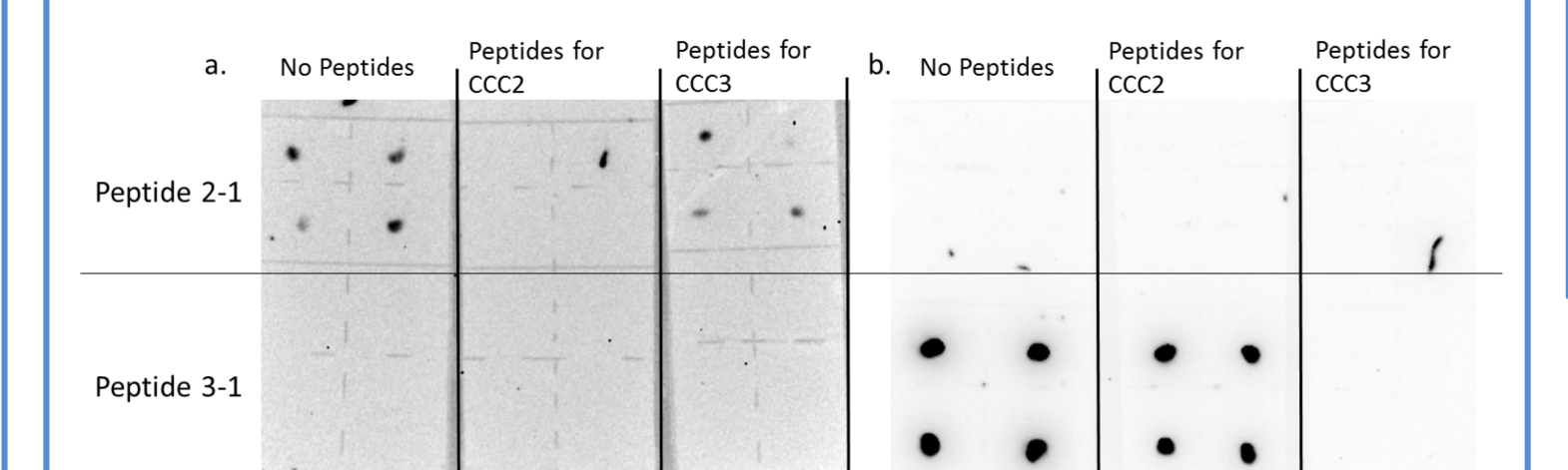


# Methods

**Mosquito Rearing** *A. aegypti* larvae and adults were reared and maintained as described (Pannabecker et al., 1993). Eggs (Liverpool) were hatched under a vacuum filter to induce low  $O_2$ . Larvae were raised in a container filled with fresh water (1.5 in) in a controlled environment at 28°C and a light:dark cycle of 14:10 hr. Larval food consists of 3 parts finely ground TetraFin fish flakes (Tetra, Melle, Germany) and 1 part Yeast Extract (Spectrum Chemical). The adults mosquitoes were fed 10% sucrose/tap water using an inverted test-tube plugged with cotton.

**Tissue Collection** Sugar-fed mosquitoes and 4th instar larval renal organs (Malpighian Tubules, hindgut, and anal papillae) were dissected as described (Pannabecker et al., 1993). Whole body samples were immediately placed in a collection tube containing 50µL of 1% Triton-X 100 and 1% SDS in Ringer Solution (in mM: NaC1 150, HEPES 25, KC13.4, CaC12 1.7, NaHCO3 1.8, MgC12 0.6, and glucose 5). Protein concentrations were determined with the bicinchoninic acid method (BCA Reagent, Pierce Chemical).

**Figure 4.** Western blots of whole body samples from adult female mosquitoes (W.F.), 4th instar larvae (W.L), adult males (W.M.), and whole pupae (W.P.) Image a. and c. come from the same rabbit and b. and d. come from the other. The images were exposed for A) 6 seconds, B) 15 seconds, C) 50 seconds, D) 50 seconds.



250 kD 250 kD 250 kD 250 kD 250 kD 150 kD 15

**Figure 6.** Peptide competition Western Blots of adult female mosquitoes (W.F.) and 4th instar larvae (W.L). Image Blots were washed with an unpurified antiserum When blocking, the peptides were added to the antibody such that the antibodies would bind to the free peptide before binding to the PVDF film. The images were exposed at 45 sec.

## Reference

 Gillen CM, Blair CR, Heilman NR, Somple M, Stulberg M, Thombre R et al (2006) The cation-chloride cotransporter, masBSC, is widely expressed in Manduca sexta tissues. J Insect Physiol 52:661–668
Rodan, A.R., Baum, M., Huang, C.-L., 2012. The Drosophila NKCC Ncc69 is required for normal renal tubule function. Am. J. Physiol. Cell Physiol. 303, C883-C894.

3. Pannabecker TL, Hayes TK, and Beyenbach KW., 1993. Regulation of epithelial shunt conductance by the peptide leucokinin. J. Membr. Biol. 132: 63–76.

Acknowledgement

Western Blot Proteins were separated on 7.5% Tris–HCI gels and transferred to PVDF membranes. Blots were blocked for 1 h in phosphate buffered saline with 0.1% Tween 20 (PBT) 5% bovine serum albumin (BSA), and exposed to primary antibody for 3 hr with gentle agitation. Unpurified antiserum was used at dilutions ranging from 1:1000 to 1:5000. Purified aeCCC2 and aeCCC3 antibody was used at a 1:1000 and 1:5000 dilution, respectively, of a 1 mg/ml stock solution. Blots were washed 3 times for 5 min in PBT 5% BSA, 3 times for 5 min in PBT, exposed to HRP-conjugated anti-rabbit IgG secondary antibody (Sigma Chemical, 1:5000 dilution) and the washes were repeated. Detection was found by using chemiluminescence (ChemiDoc MP, Bio Rad).

**Figure 5.** Peptide competition Dot Blots. Image a. is the aeCCC2 antibody while image b. is the aeCCC3 antibody. Peptides used in producing the aeCCC2 and aeCCC3 antibodies were spotted onto PVDF film. When blocking, the peptides were added to the purified antibody such that the antibodies would bind to the free peptide before binding to the PVDF film. The images were exposed at 70 sec.

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